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1	UNITED STATI	ES DISTRICT COURT	
2	NORTHERN DISTRICT OF CALIFORNIA		
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4	THE SAC AND FOX NATION OF OKLAHOMA	Case No. 21-md-02996-CRB (SK)	
5		ORIGINAL COMPLAINT	
6	Plaintiff,	REDACTED FOR PUBLIC FILING	
7	VS.	JURY TRIAL DEMANDED	
8	MCKINSEY & CO., INC.; MCKINSEY HOLDINGS, INC.; MCKINSEY &		
9	COMPANY, INC. UNITED STATES; MCKINSEY & COMPANY, INC.		
10	WASHINGTON D.C.; PUBLICIS HEALTH, LLC; PRACTICE FUSION, INC.; ALLSCRIPTS HEALTHCARE		
11	SOLUTIONS, INC.; and ZS ASSOCIATES, INC.		
12	Defendants.		
13	Dejenaunis.		
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I. INTRODUCTION

- 1. For two decades now, the modern¹ opioid crisis has raged. In 2017, the President of the United States announced a public health emergency. In 2020, drug-overdose deaths in the United States soared nearly 30%.² Drug overdoses now kill more than 100,000 Americans per year more than vehicle crash and gun deaths combined.³ Today, there are fewer and fewer Americans whose lives have not been scarred by the epidemic.
- 2. History repeats itself not through accident or accretion, but through affirmative acts undertaken by those who wish it so. The modern epidemic has a cause, and this complaint seeks to hold accountable principal participants in its creation and propagation. The modern epidemic began with the introduction and expansive promotion of OxyContin by Purdue Pharma in 1996 and was driven thereafter by industry-wide marketing efforts in which Defendants were principal and knowing participants. It continues to rage today, further enflamed by the isolation of the COVID-19 pandemic.⁴
- 3. The pharmaceutical industry is complex and highly regulated. Drug manufacturers cannot, and do not, do everything on their own. These innovative companies endeavor to research,

¹ Throughout history, instances of widespread availability of opioids causing severe detriment to all of us are legion. To begin with, the drug was powerful enough to be used as a means of colonial expansion by the British Empire in China. There were two Opium Wars. See W. Travis Hanes III & Frank Sanello, The Opium Wars: The Addiction of One Empire and the Corruption of Another, 2002; Julia Lovell, The Opium War: Drugs, Dreams, and the Making of Modern China, 2011.

More recently, the United States has experienced previous waves of opioid abuse. "The first was in the early 1900's, when heroin was marketed alongside Bayer aspirin as a remedy for numerous minor ailments." Dr. Anna Lembke, *Drug Dealer, M.D.*, Johns Hopkins University Press (2016), Pg. 57. A second heroin epidemic struck the United States during the Vietnam War. *Id. See also* John Fauber et al., "A Look Back: Abandoned Painkiller Makes a Comeback," *Medpage Today*, June 9, 2017, *available at*: https://www.medpagetoday.com/psychiatry/addictions/65916 (describing the removal of Numorphan, an oxymorphone product, from the market in the 1970's in response to widespread abuse). Throughout the time-period relevant to this complaint, the dangers of widespread opioid availability weren't just common knowledge, they were historical fact.

² Betsy McKay, "U.S. Drug-Overdose Deaths Soared Nearly 30% in 2020, Driven by Synthetic Opioids," *Wall Street Journal*, July 14, 2020, *available at*: https://www.wsj.com/articles/u-s-drug-overdose-deaths-soared-nearly-30-in-2020-11626271200

³ German Lopez, "A Rising Death Toll," New York Times, February 13, 2022, available at: https://www.nytimes.com/2022/02/13/briefing/opioids-drug-overdose-death-toll.html.

⁴ See https://healthmatters.nyp.org/an-overdose-epidemic-amid-the-pandemic/

develop, obtain approval, and bring to market products that improve the health and livelihood of hundreds of millions of people worldwide. Because the scale of the industry is vast, the stakes – both in terms of potential profit and impact on human lives – are high, and the industry's complexity byzantine. From cutting-edge medical and scientific research, to navigating the state and federal regulatory environment, to marketing their drugs in a responsible manner to prescribers and the consuming public, the demands on a traditional pharmaceutical manufacturer are multi-faceted, ever-present, and continuously changing.

- 4. The reality is pharmaceutical manufacturers routinely rely on third parties to design, implement, and oversee projects and workflows to achieve mission-critical tasks. Given the complexity of the industry, there are numerous companies that find their niche in offering core services to pharmaceutical manufacturers that are critical to the success of the manufacturers' operations but are not performed by the manufacturer alone. These third parties are necessary components of the drug manufacturing and sales industry as a whole.
- 5. Manufacturers do not rely on these third-party providers on a one-off basis, but instead rely on companies like McKinsey, Publicis, ZS, and Practice Fusion again and again to design and implement measures to achieve specific needs. The relationships are recurring and long-term. It is not uncommon for McKinsey or ZS or Publicis to advise multiple pharmaceutical manufacturers regarding the sales and marketing of competing products, such as branded extended-release opioids. Further, it is common for third-party consultants such McKinsey, ZS and Publicis to commoditize their business intelligence in a way which expands markets across the industry. In other words, they often sell the same strategy to multiple clients. This complaint concerns the conduct of these providers.
- 6. One Defendant McKinsey is the preeminent management consultancy on Earth.

 Another Publicis is one of the largest advertising agencies on Earth. Publicis Groupe S.A. is the

French parent of Defendant Publicis Health LLC ("Publicis"). A third Defendant is a little known, but principal architect of the sales and marketing efforts that begat the opioid crisis, pharmaceutical consulting firm ZS Associates, Inc. ("ZS"). A fourth, Practice Fusion, Inc. ("Practice Fusion") offered a unique channel through which the marketing messages and strategy created by co-Defendants could be delivered to the intended audience: healthcare providers, the folks with prescription pads in their hands. All were crucial components of and contributors to the architecture and functioning of the opioid marketplace.⁵

- 7. McKinsey is a management consulting firm with operations across the globe. It played a central role in the unfolding, propagation, and exploitation of the opioid crisis by advising multiple opioid manufacturers and other industry participants how to sell as many opioids as conceivably possible. Knowing that its clients' products were highly addictive, ineffective, and unsafe for the treatment of long-term chronic pain, non-acute pain, and non-cancer pain, McKinsey developed a singular focus on increasing opioid sales, no matter the resultant cost to society. McKinsey did this for well over a decade, despite knowing full well the risk to public health and safety and the widespread economic harm from developing and implementing the transformation of strictly controlled substances into top-selling blockbuster drugs.
- 8. The purpose of McKinsey's work with its opioids clients was at all times to maximize return on investment. The whole point for those clients (and hence McKinsey) was to make as much money as possible. They all did. This relentless drive to increase sales and create greater availability of opioids was made with no concern about the parallel, known, and inevitable increase in opioid-related deaths, addiction, abuse, diversion, and misuse.
- 9. In the world of management consulting, McKinsey is preeminent. It is one of the world's oldest, largest, and most lucrative consulting firms and is generally seen as the most

⁵ The fifth defendant, Allscripts, Inc. ("Allscripts") is the parent company of defendant Practice Fusion.

prestigious firm in the industry. More consiglieri than one-off advisor, McKinsey touts its model of engaging in "transformational partnerships" with its clients. McKinsey learns each client's business intimately, embeds itself into all levels of the corporate hierarchy, and provides granular strategies to achieve transformative goals for its clients.

- 10. Marvin Bower, the managing director of McKinsey from 1950 to 1967, was "the father of the consulting profession." He "turned the business of selling management advice into a keystone of American corporate culture," and is "credited with taking a fledgling industry and setting its course not only as to the kinds of services it could sell but also the standards it must uphold for its work to be respected." A lawyer by trade, Bower stressed that management consulting should be seen as an emergent profession, akin to the law or accounting, with obligations to clients and to the broader society that extend beyond the mere commercial.
- 11. Bower instilled an ethos at McKinsey that has been reinforced throughout the decades as a core value of the firm: "Deliver the bad news if you must, but deliver it properly."8 Bower's principles, and the values he imparted within McKinsey, are said to guide the firm to the present day. "In many ways, certainly in spirit and soul, Marvin continued to lead it after he retired, and he leads it still," eulogized Rajat Gupta, McKinsey's then-global managing partner, at Bower's funeral in 2003.9

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⁶ Douglas Martin, Marvin Bower, 99; Built McKinsey & Co., N.Y. Times, Jan. 24, 2003, available at: https://www.nytimes.com/2003/01/24/business/marvin-bower-99-built-mckinsey-co.html ⁷ *Id*.

⁸ Duff McDonald, *The Firm* 35 (2014).

⁹ Id. at 270. In many ways, Gupta was an interesting figure to opine on Bower's legacy. Indeed, Gupta's leadership of McKinsey is in many respects to be contrasted with Bower's legacy. Many of the values Bower emphasized—an emphasis on professionalism over commercial exploitation, for example—were jettisoned under Gupta's tenure as managing partner of the firm, which ended in 2003. "Under his watch, McKinsey began to chase top billings in a way it never had before." Id. at 234. For instance, McKinsey first began accepting equity stakes in clients as a form of incentive compensation during Gupta's tenure. Previously, McKinsey only charged standard fees for its consulting services as Bower disdained the notion of taking equity stakes in clients. Id. at 234. Under Gupta, McKinsey also began to allow consultants' compensation to be tied to client performance. *Id*.

Consistent with Gupta's efforts to monetize McKinsey's consulting business in ways previous firm leadership had not, McKinsey also began to expand its client base. "While the firm would never admit as much, under Gupta, McKinsey began working for just about anyone with a fat bank account and a checkbook." Id. at 266.

- 12. This case is, in large part, about the firm's failure to adhere to Bower's simple, foundational tenet. It arises instead from the firm's steadfast and continual work to maximize opioid sales in partnership with numerous clients during the pendency of the worst man-made epidemic in modern medical history. It is about McKinsey never delivering the "bad news" of opioids' devastating impact on Plaintiff and the public, and instead looking the other way for money.
- 13. When it came to opioids, McKinsey did far more than just give advice. Not only did it suggest courses of action that its clients should adopt, the firm remained in place and worked collaboratively alongside its clients to actually implement McKinsey's recommendations to achieve objectives jointly identified by the clients and McKinsey. McKinsey stood alongside its clients in the arena doing the deeds.
- 14. The deceptive marketing strategies that McKinsey and its clients invented, developed, deployed, and continually refined for years to expand the market for opioids are foundational to the epidemic.
- 15. McKinsey worked hand-in-hand with major opioid manufacturers, including Purdue Pharma L.P., Endo Pharmaceuticals, ¹⁰ Johnson & Johnson, ¹¹ and Mallinckrodt ¹² for years. At the same time, McKinsey advised other participants in the opioid supply chain, including distributors, pharmacies, and even regulators.

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Institutions age, and by the time Gupta came to lead the firm in 1994, McKinsey was a mature institution. It had built up significant value in its reputation by historically advising only "blue chip" companies "at the top of the corporate pyramid." Id. Under Gupta, McKinsey began the process of realizing that value. For McKinsey, the way to monetize an elite reputation was to start advising those it historically may have shunned as clients—to start offering its imprimatur, in addition to its services, for money. McKinsey's work with opioid manufacturers began under Gupta's leadership.

¹⁰ "Endo Pharmaceuticals" or "Endo" refers to Endo Health Solutions Inc., Endo International plc, and Endo Pharmaceuticals Inc., collectively.

¹¹ "Johnson & Johnson" refers to Johnson & Johnson Services, Inc. and its wholly-owned subsidiary Janssen Pharmaceuticals, Inc. ("Janssen").

¹² "Mallinckrodt" refers to Mallinckrodt LLC and Mallinckrodt plc, together.

16. In particular, McKinsey advised the Sackler family and their company, Purdue, for years while Purdue aggressively marketed OxyContin, widely viewed as the taproot of the opioid crisis. The relationship began no later than 2004. In the years following Purdue's 2007 guilty plea for misleadingly marketing OxyContin, McKinsey continued to work closely with Purdue to dramatically increase OxyContin sales, notwithstanding the existence of a five-year Corporate Integrity Agreement that Purdue entered as part of its guilty plea.

- 17. McKinsey knew of the dangers of opioids and in particular the prior misconduct of Purdue but nonetheless advised Purdue and other opioid manufacturers to improperly market and sell OxyContin and other prescription opioids, supplying granular sales and marketing strategies and remaining intimately involved throughout implementation of those strategies. McKinsey's actions resulted in a surge in sales of OxyContin and other opioids that fueled and prolonged the opioid crisis.
- 18. For years, McKinsey advised Purdue on, designed, and helped to implement various strategies to raise sales of OxyContin by focusing on high dose sales and deceptively messaging to physicians that OxyContin would improve function and quality of life. For example, McKinsey urged Purdue to maximize sales by dictating, to a greater degree, which prescribers its sales representatives would target, exploring ways to increase the amount of time those sales representatives spent in the field increasing opioid sales and prioritizing OxyContin in incentive compensation targets.¹³
- 19. McKinsey's partnership with Purdue reached its fever pitch in the summer of 2013. In January of that year, Purdue's Corporate Integrity Agreement expired, and Purdue was no longer bound by its constraints. Within months, the Sacklers tasked McKinsey with transforming Purdue's

¹³ PPLPC012000437346

approach to OxyContin sales in order to extract as much money as possible from the remaining patent life of the drug.¹⁴

- 20. In response, McKinsey developed and proposed Project Turbocharge, a series of transformational changes that McKinsey proposed to implement at Purdue to dramatically increase OxyContin sales by re-tooling Purdue's sales force and investing large amounts of capital to "turbocharge" it. "[O]ur recommendation is that Purdue makes a clear go-no-go decision to 'Turbocharge the Sales Engine'," McKinsey told Purdue on August 8, 2013.
- 21. The Sacklers chose "go," and McKinsey subsequently implemented and continually refined Project Turbocharge at Purdue over the course of years, to devastating, but profitable, effect.
- 22. McKinsey has recently been the subject of scrutiny for its various business practices, including its work facilitating the opioid crisis with Purdue.¹⁵ On March 7, 2019, Kevin Sneader, McKinsey's then-global managing partner, addressed all McKinsey employees regarding this scrutiny. Drawing inspiration from Theodore Roosevelt, Sneader stated,

[W]e cannot return to a time when we were in the background and unobserved. Those days have gone. Indeed, I have little doubt that scrutiny—fair and unfair—will continue. It is the price we pay for being "in the arena" and working on what matters.¹⁶

As it happens, Mr. Sneader is not the only McKinsey person to draw inspiration from Roosevelt. *Citizenship in a Republic* similarly inspired Dominic Barton, the man Mr. Sneader succeeded as McKinsey's global managing

¹⁴ OxyContin, like any branded pharmaceutical, is subject to eventual patent expiration and competition from generic opioid manufacturers.

¹⁵ See Michael Forsythe and Walt Bogdanich, McKinsey Advised Purdue Pharma How to 'Turbocharge' Opioid Sales, Lawsuit Says, N.Y. Times, Feb. 1, 2019, available at: https://www.nytimes.com/2019/02/01/business/purdue-pharma-mckinsey-oxycontin-opiods.html.

¹⁶ See "The Price We Pay for Being 'In the Arena'": McKinsey's Chief Writes to Staff About Media Scrutiny and Scandal, Fortune Magazine, March 8, 2019, available at https://fortune.com/2019/03/08/mckinsey-staff-letter-kevinsneader/. The "arena" reference is to Citizenship in a Republic, a speech delivered by Theodore Roosevelt at the Sorbonne on April 23, 1910:

It is not the critic who counts; not the man who points out how the strong man stumbles, or where the doers of deeds could have done them better. The credit belongs to the man who is actually in the arena [here, McKinsey; and the arena, opioid sales], whose face is marred by dust and sweat and blood; who strives valiantly; who errs, who comes short again and again, because there is no effort without error and shortcoming; but who does actually strive to do the deeds; who knows great enthusiasms, the great devotions; who spends himself in a worthy cause; who at the best knows in the end the triumph of high achievement, and who at the worst, if he fails, at least fails while daring greatly, so that his place shall never be with those cold and timid souls who neither know victory nor defeat.

- 23. Weeks later, McKinsey announced that it would no longer work for any opioid manufacturer. "Opioid abuse and addiction are having a tragic and devastating impact on our communities. We are no longer advising clients on any opioid-specific business and are continuing to support key stakeholders working to combat the crisis."¹⁷
- 24. The price for being in the arena is more than mere scrutiny. McKinsey is liable for its misconduct and the harms it caused or exacerbated. McKinsey is liable for its successful efforts to increase opioid sales for years. It continued this work unabated and with alacrity despite events as stunning as Purdue's 2007 guilty plea for misbranding OxyContin, Purdue's 2015 settlement with the State of Kentucky, and numerous other enforcement actions related to opioid sales and marketing by McKinsey clients. Through it all, McKinsey remained steadfast in its efforts to promote opioid sales for all of its clients for the purpose of maximizing return on investment without regard to the obvious implications of what they were doing. Indeed, the firm endeavored alongside its clients to increase the size of the overall opioid market for nearly two decades, until as late as March 22, 2019, despite increasingly blood-red flags along the way. 18
- 25. And McKinsey was not alone; Defendants Publicis, ZS, and Practice Fusion endeavored alongside McKinsey, shoulder to shoulder and in common cause with McKinsey and their manufacturer clients to perpetuate and increase in size the opioids market; to sell more and more pills.

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partner. It served as the basis for his 2017 address to the Ivey Business School in Canada. See Dominic Barton In the Arena: Leadership in an Age of Disruption, October 17, 2017, available at: https://www.ivey.uwo.ca/media/3780710/ daquino_lecture2017.pdf. While McKinsey continues to preach the values of corporate integrity from the Bower area, its actions show that that professed moral compass needs to be serviced; it doesn't work anymore.

¹⁷ See Paul La Monica, Consulting firm McKinsey no longer working with opioid maker Purdue Pharma, CNN, May 24, 2019, available at: https://www.cnn.com/2019/05/24/business/mckinsey-purdue-pharma-oxycontin/index.html. The statement was attributed to McKinsey as an entity. No individual's name was attributed.

¹⁸ See "About McKinsey's past work for opioid manufacturers," last updated March 22, 2021, available at: https://www.mckinseyopioidfacts.com ("We decided nearly two years ago to end all work on opioid-specific business ")

- 26. Publicis is one of the "Big Four," the four firms that account for more than half of the global advertising industry. In 2002, as the opioid crisis was taking root across the United States, the president of the American Association of Advertising Agencies, stated, "Now you have four megacompanies with revenues that are staggering, bigger than some of the companies they serve." The rise of the Big Four came through decades of mergers and acquisitions of separate agencies; each is essentially a conglomerate. By October of 2021, Publicis had risen to become the largest advertising conglomerate in the world, with a market capitalization in excess of \$16 billion.²⁰
- 27. Publicis has acquired and developed particular expertise in serving the pharmaceutical industry. Publicis through its's myriad divisions serves multiple pharmaceutical manufacturers in advertising their drugs. It relies on the healthcare sector for approximately 13% of its annual revenue, making it one of the largest industry sectors by revenue for the conglomerate.²¹
- 28. The over-marketing of opioids schedule II controlled substances that are *controlled* because they are known to be addictive and deadly was the wellspring of our national crisis. Although the introduction of OxyContin by Purdue Pharma L.P. ("Purdue") in the late 1990's is widely acknowledged as a precipitating cause of the opioid crisis, Purdue was not the only pharmaceutical company to enthusiastically foment and exploit the booming market of controlled substances used for the treatment of pain. An industry-wide sales and marketing effort was deployed over the years by numerous manufacturers of opioid medications in order to maximize the amount of opioids they could sell.
- 29. The sales and marketing efforts to sell opioids to as many individuals as possible, even when they were known to be addictive, were not solely designed by the manufacturers

¹⁹ Stuart Elliott, Advertising's Big Four: It's Their World Now, *New York Times*, March 31, 2002, *available at*: https://www.nytimes.com/2002/03/31/business/advertising-s-big-four-it-s-their-world-now.html

²⁰ See https://www.prweek.com/article/1731568/publicis-overtakes-rivals-worlds-valuable-agency-group

²¹ See http://documents.publicisgroupe.com/resultat2021/Presentation-H1-2021-RESULTS.pdf at 15.

themselves, nor did the manufacturers implement these tactics on their own. Rather, pharmaceutical manufacturers routinely relied on Defendants to design and implement crucial aspects of the sales and marketing strategies used to sell opioids.

- 30. Publicis did not produce mere copy. Publicis not only designed these sales and marketing campaigns for numerous opioid manufacturers, it also worked in identifying the optimal targets for the different messages Publicis delivered on behalf of its clients. In many instances, pharmaceutical manufacturers would outsource practically the entire business of selling its drugs. Through Publicis Touchpoint Solutions, Publicis provided entire sales forces on a contract basis to be used by their manufacturer clients to detail prescribers.
- 31. Nor was Publicis merely some ancillary vendor. Because of its partnerships with multiple clients selling competing branded opioid products contemporaneously, Publicis's role was unique. It served as a hub, aggregating knowledge of what numerous competitors within the industry were doing with respect to designing sales and marketing campaigns, contemporaneously. Indeed, Publicis worked with industry-wide groups to address challenges that the entire opioids industry faced and compiled invaluable insights and business intelligence which it commoditized by providing it to numerous clients.
- 32. As a hub, Publicis also connected opioid manufacturers to specialist vendors, such as Defendant Practice Fusion, who worked with Publicis and Purdue to target and deliver content to healthcare providers designed to increase the amount of Purdue's opioids sold through Practice Fusion's proprietary software platform used in doctors' offices across the country.
- 33. ZS is a private consulting company that specializes in the pharmaceutical industry. It was founded in 1983 by two professors at the eminent Kellogg School of Business at Northwestern University. Since then, ZS has achieved substantial growth, and now employs thousands of consultants and enjoys hundreds of millions of dollars in annual revenue. "ZS" is the

initials of the two founders, Professors Andris Zoltners and Prabhakant Sinha. In 2020 ZS had 8,000 employees and was listed as one the Best Management Consulting Firms by Forbes in 2021.²²

- 34. In particular, ZS specializes in providing critical pharmaceutical sales and marketing services to drive increased sales volume and related profits.
- 35. As set forth in this complaint, Defendants' purpose in working with these companies was plain and singular: to maximize profits for their clients by making sure that every dollar spent on sales and marketing of opioids generated as many sales of these addictive controlled substances as possible. Maximizing profits and revenue for Defendants' clients was achieved by maximizing the total volume of opioids sold. McKinsey, Publicis and ZS applied their sales and marketing acumen to multiple opioid brands on behalf of numerous manufacturer clients, and often at the same time. McKinsey, Publicis and ZS were common denominators throughout.
- 36. These Defendants played a central role in the creation, prolongation, and exploitation of the opioid crisis for money. As the alarm bells sounded repeatedly in the early years of the unfolding crisis, Defendants like McKinsey continued their work with opioid manufacturers unabated and with alacrity, and for decades, as the bells rose to cacophony. It continued right up until the bitter end.²³
- 37. Moreover, the work was conducted in the shadows. Defendants treat their client relationships as confidential. Classically, ZS or Publicis like McKinsey works behind the scenes and does not publicize its work. Until recent consultant litigation, the public had essentially no knowledge or awareness of the extent of involvement that Defendants like McKinsey had in tearing apart our social fabric for profit. The Sac and Fox Nation, with its eyes now fully open to

²² See https://www.forbes.com/companies/zs/?sh=12e5e93027d0

²³ For instance, ZS had an active contract with Purdue Pharma L.P. ("Purdue") to assist with the sales and marketing of OxyContin in 2018, after Purdue had already pled guilty, settled numerous prior lawsuits brought by the DOJ, state Attorneys General and individuals and faced a new wave of similar litigation. That year, with its contract with ZS still active, Purdue chose to disband its sales force and cease marketing the drug altogether.

the true scope of the origins and prolongation of the opioid crisis, seek to hold all those responsible accountable.

II. JURISDICTION AND VENUE

- 38. Plaintiff brings this action by filing directly in in the Northern District of California pursuant to Paragraph 10 of this Court's Case Management Order Dated November 20, 2121 (Doc. #293). Plaintiff reserves the right to have this matter transferred to the United States District Court in which it would have originally filed this case absent consolidation, the United States District Court for the Western District of Oklahoma.
- 39. The United States District Court for the Western District of Oklahoma has subject matter jurisdiction over this action pursuant to 28 U.S.C. § 1332(d)(2), because (i) at least one member of the putative Class is a citizen of a state different from Defendants (ii) the amount in controversy exceeds \$5,000,000, exclusive of interest and costs, and (iii) none of the exceptions under the subsection apply to this action.
- 40. The United States District Court for the Western District of Oklahoma has personal jurisdiction over the Defendants because Plaintiffs claims arise out of, or relate to, Defendants' contacts with Oklahoma.
- 41. The United States District Court for the Western District of Oklahoma has personal jurisdiction over the Defendants because Plaintiffs claims arise out of, or relate to, Defendant's contacts with Oklahoma.
- 42. At all times relevant hereto, Defendants engaged in the business of researching, designing, and implementing sales and marketing strategies for various opioid manufacturers including Purdue Pharma in the State of Oklahoma and Caddo County and the territorial limits of the Sac and Fox Nation of Oklahoma.

- 43. The United States District Court for the Western District of Oklahoma has jurisdiction over Defendants due to Defendant's conduct in Lincoln County and throughout Oklahoma. Defendants have deliberately engaged in significant acts and omissions within State of Oklahoma and Lincoln County and the territorial limits of the Sac and Fox Nation of Oklahoma that have injured Plaintiff's residents. Defendants purposefully directed their activities in State of Oklahoma and Lincoln County and the territorial limits of the Sac and Fox Nation of Oklahoma and its residents, and the claims arise out of those activities.
- 44. Venue is proper in the United States District Court for the Western District of Oklahoma because a substantial part of the events giving rise to Plaintiff's claims occurred in, were directed to, and/or emanated from this District. 28 U.S.C. § 1391(b).

III. PARTIES

- 45. Plaintiff Sac and Fox Nation of Oklahoma is a sovereign Indian Tribe recognized by federal, state, and tribal law. The Sac and Fox Nation's jurisdictional territory encompasses Lincoln County, Oklahoma. Plaintiff is not a citizen of any state for purposes of diversity jurisdiction.
- 46. The Sac and Fox Nation exercises inherent and constitutional governmental authority on behalf of the Nation itself and its members. The Sac and Fox Nation brings this action on its own behalf and on behalf of its members and citizens in the public interest to protect the health, safety, and welfare of the citizens of the Sac and Fox Nation. The Sac and Fox Nation does so in an effort to address the opioid addiction epidemic within the Sac and Fox Nation and to recover damages and seek other redress for the harms cause by Defendant's conduct.
- 47. Defendant McKinsey & Company, Inc. is a corporation organized under the laws of the state of New York. McKinsey's principal place of business is located at 711 Third Avenue,

New York, NY 10017. It may be served with process via its registered agent, Corporation Service Company, at 80 State Street, Albany, NY 12207.

- 48. Defendant McKinsey Holdings, Inc. is a Delaware corporation with its principal place of business is located at 711 Third Avenue, New York, NY 10017. It may be served with process via its registered agent, Corporation Service Company, 251 Little Falls Drive, Wilmington, DE 19808
- 49. Defendant McKinsey & Company, Inc. United States is a Delaware corporation with its principal place of business is located at 711 Third Avenue, New York, NY 10017. It may be served with process via its registered agent, Corporation Service Company, 251 Little Falls Drive, Wilmington, DE 19808
- 50. Defendant McKinsey & Company, Inc. Washington D.C. is a Delaware corporation with its principal place of business is located at 711 Third Avenue, New York, NY 10017. It may be served with process via its registered agent, Corporation Service Company, 251 Little Falls Drive, Wilmington, DE 19808.
- 51. Upon information and belief, McKinsey & Company, Inc. is the parent company of McKinsey & Company Holdings, Inc., which is itself the parent company of both McKinsey & Company, Inc. United States and McKinsey & Company, Inc. Washington D.C. Upon information and belief, each subsidiary corporation is wholly-owned by its parent. Despite the corporate form, McKinsey began as a partnership and still refers to its senior employees as "partners." Those partners are the firm's shareholders. Collectively, these four Defendants are referenced throughout as "McKinsey."
- 52. McKinsey a is global management consultancy with offices in over 130 cities in 65 countries, including the following United States cities: Atlanta, GA; Austin, TX; Houston, TX; Dallas, TX; San Francisco, CA; Los Angeles, CA; Redwood City, CA; Boston, MA; Charlotte,

NC; Chicago, IL; Cleveland, OH; Denver, CO; Detroit, MI; Miami, FL; Miramar, FL; Tampa, FL; Minneapolis, MN; Summit, NJ; New York, NY; Philadelphia, PA; Pittsburgh, PA; Seattle, WA; St. Louis, MO; Stamford, CT; Waltham, MA; and Washington, D.C.

- 53. McKinsey is registered to do business in all fifty states.
- 54. Defendant Publicis Health, LLC ("Publicis Health") is a Delaware limited liability company with its principal place of business in New York City. Additionally, Publicis maintains an office within the State at 100 E Penn Square, 11th Floor, Philadelphia, PA 19107, and is registered to do business in Pennsylvania. It may be served with process through its registered agent, Corporation Service Company, 2595 Interstate Dr. #103, Harrisburg, PA 17110.
- 55. Defendant Practice Fusion, Inc. ("Practice Fusion") is a Delaware corporation with its headquarters in San Francisco, California. It may be served with process through its registered agent, National Registered Agents, Inc. located at 818 West Seventh St., Ste 930, Los Angeles, CA 90017. Practice Fusion provided electronic health records ("EHR") services to clinicians and healthcare providers, and was ultimately acquired by Defendant Allscripts.
- 56. Defendant Allscripts Healthcare Solutions, Inc. ("Allscripts") is a Delaware corporation with its headquarters in Chicago, Illinois. On February 13, 2018, Allscripts completed a merger whereby it acquired Defendant Practice Fusion and became a successor in interest thereto.
- 57. Defendant ZS Associates, Inc., is a foreign corporation with its principal office located at 1800 Sherman Avenue, Evanston, Illinois 60201. It may be served with process through its registered agent, Illinois Service Corporation, 801 Adlai Stevenson Dr., Springfield, IL 62703.

IV. FACTUAL ALLEGATIONS

58. Organizationally, this complaint will tell the individual stories of each defendant, beginning with McKinsey, then Publicis, then Practice Fusion, and finally ZS, and will detail each defendant's interactions with multiple opioid manufacturers – and each other – in turn. As will be

seen, each worked for the same opioid manufacturers at the same time and on the same projects, and all were unified in their common purpose²⁴: to maximize opioid sales and associated profits for the past two decades.

a. The Opioid Crisis

- 59. The term "opioid" refers to a class of drugs that bind with opioid receptors in the brain and includes natural, synthetic, and semi-synthetic opioids. Natural opioids are derived from the opium poppy. Generally used to treat pain, opioids produce multiple effects on the human body, the most significant of which are analgesia, euphoria, and respiratory depression.
- 60. The opium poppy contains various opium alkaloids, three of which are used in the pharmaceutical industry today: morphine, codeine, and thebaine. Early use of opium in Western medicine was a tincture of opium and alcohol called laudanum, which contains all of the opium alkaloids and is still available by prescription today. Chemists first isolated the morphine and codeine alkaloids in the early 1800s.
- 61. In 1827, the pharmaceutical company Merck began large-scale production and commercial marketing of morphine. During the American Civil War, field medics commonly used morphine, laudanum, and opium pills to treat the wounded, and many veterans were left with morphine addictions. By 1900, an estimated 300,000 people were addicted to opioids in the United States, and many doctors prescribed opioids solely to prevent their patients from suffering withdrawal symptoms. The nation's first Opium Commissioner, Hamilton Wright, remarked in 1911: "The habit has this nation in its grip to an astonishing extent . . . Our prisons and our hospitals are full of victims of it, it has robbed ten thousand businessmen of moral sense and made them

²⁴ See "McKinsey on Implementation", McKinsey & Company Inc., April 30, 2017, available at: https://www.youtube.com/watch?v=rEQOGVpl9CY ("One thing intriguing about the engagements is that they often have a common purpose, and a genuine cause," Josh, a Senior Implementation Coach at McKinsey, explained.)

beasts who prey upon their fellows . . . it has become one of the most fertile causes of unhappiness and sin in the United States."

- 62. Pharmaceutical companies have long tried to develop substitutes for opium and morphine that would provide the same analgesic effects without the addictive properties. In 1898, Bayer Pharmaceutical Company began marketing diacetylmorphine (obtained from acetylation of morphine) under the trade name "Heroin." Bayer advertised heroin as a non-addictive cough and cold remedy suitable for children, but as its addictive nature became clear, heroin distribution in the United States was limited to prescription only in 1914 and then banned altogether a decade later.
- 63. Although heroin and opium became classified as illicit drugs, there is little difference between them and prescription opioids. Prescription opioids are synthesized from the same plant as heroin, have similar molecular structures, and bind to the same receptors in the human brain.
- 64. Due to concerns about their addictive properties, prescription opioids have usually been regulated at the federal level as Schedule II controlled substances by the Drug Enforcement Administration since 1970.
- 65. Throughout the twentieth century, pharmaceutical companies continued to develop prescription opioids like Percodan, Percocet, and Vicodin, but these opioids were generally produced in combination with other drugs, with relatively low opioid content.
- 66. In contrast, OxyContin, the product whose launch in 1996 ushered in the modern opioid epidemic, is pure oxycodone. Purdue initially made it available in the following strengths: 10 mg, 15 mg, 20 mg, 30 mg, 40 mg, 60 mg, 80 mg, and 160 mg. The weakest OxyContin delivers as much narcotic as the strongest Percocet, and some OxyContin tablets delivered sixteen times that.

- OxyContin and MS Contin, Janssen's Nucynta ER and Duragesic, Endo's Opana ER, and Actavis's Kadian, are designed to be taken once or twice daily and are purported to provide continuous opioid therapy for, in general, twelve hours. Short-acting opioids, such as Cephalon's Actiq and Fentora, are designed to be taken in addition to long-acting opioids to address "episodic pain" (also referred to as "breakthrough pain") and provide fast-acting, supplemental opioid therapy lasting approximately four to six hours. Still other short-term opioids, such as Insys's Subsys, are designed to be taken in addition to long-acting opioids to specifically address breakthrough cancer pain, excruciating pain suffered by some patients with end-stage cancer. The opioid manufacturers promoted the idea that pain should be treated by taking long-acting opioids continuously and supplementing them by also taking short-acting, rapid-onset opioids for episodic or "breakthrough" pain.
- 68. Patients develop tolerance to the analgesic effect of opioids relatively quickly. As tolerance increases, a patient typically requires progressively higher doses in order to obtain the same perceived level of pain reduction. The same is true of the euphoric effects of opioids—the "high." However, opioids depress respiration and, at very high doses, can, and often do, arrest respiration altogether. At higher doses, the effects of withdrawal are more severe. Long-term opioid use can also cause hyperalgesia, a heightened sensitivity to pain.
- 69. Discontinuing opioids after more than just a few weeks of therapy will cause most patients to experience withdrawal symptoms. These withdrawal symptoms include severe anxiety, nausea, vomiting, headaches, agitation, insomnia, tremors, hallucinations, delirium, pain, and other serious symptoms, which may persist for months after a complete withdrawal from opioids, depending on how long the opioids were used.

70. As one doctor put it, the widespread long-term use of opioids "was an experiment on the population of the United States. It wasn't randomized, it wasn't controlled, and no data was collected until they started gathering death statistics."

71. The results were devastating, and the nation continues to reach ever grimmer milestones. In 2020, drug-overdose deaths in the United States soared nearly 30%, reaching all-time highs.²⁵

b. Selling Controlled Substances: Marketing and the Origins of the Opioid Crisis

72. Selling drugs is big business:

Perhaps the most powerful tool that pharmaceutical companies have for driving up profit margins and cultivating growth of drug markets is advertising. Pharmaceutical companies, especially the makers of opioid prescriptions, spend an enormous amount of money advertising their products – far more than they ever spend on drug research and development (Swanson 2015). In 2012 alone, the US pharmaceutical industry spent more than \$27 billion on drug promotion – including more than \$24 billion on marketing directly to physicians and \$3 billion advertising to consumers.²⁶

73. Professor Amanda Pustilnik of the Center for Law, Brain, & Behavior, and herself a former McKinsey & Company management consultant, emphasized the centrality of the role coordinated opioid sales and marketing played in creating the opioid crisis. "[T]he story of the opioid epidemic is often misrepresented as a story of irresponsible patients and over-prescribing doctors." Referring to the recent lawsuit brought by the State of Massachusetts against Publicis, Professor Pustilnik identified a more pernicious cause: the efforts by defendants to change

²⁵ Betsy McKay, "U.S. Drug-Overdose Deaths Soared Nearly 30% in 2020, Driven by Synthetic Opioids," *Wall Street Journal*, July 14, 2020, *available at*: https://www.wsj.com/articles/u-s-drug-overdose-deaths-soared-nearly-30-in-2020-11626271200.

Melina Sherman, "Opiates for the masses: constructing a market for prescription (pain)killers," Journal of Cultural Economy, Vol. 10, Issue 6, 2017, available at: https://www.tandfonline.com/doi/full/10.1080/17530350.2017.1352010 (citing Swanson, Ana, 2015. Big pharmaceutical companies are spending far more on marketing than on research. The Washington Post, February 11, available at: https://www.washingtonpost.com/news/wonk/wp/2015/02/11/big-pharmaceutical-companies-are-spending-far-more-on-marketing-than-research/">https://www.washingtonpost.com/news/wonk/wp/2015/02/11/big-pharmaceutical-companies-are-spending-far-more-on-marketing-than-research/; Cegedim Strategic Data, 2013. 2012 U.S. pharmaceutical company promotion spending. In the Pew Charitable Trust (2013), Fact Sheet: Persuading the Prescribers: Pharmaceutical Industry Marketing and its Influence on Physicians and Patients, available at: <a href="https://www.pewtrusts.org/en/research-and-analysis/fact-sheets/2013/11/11/persuading-the-prescribers-pharmaceutical-industry-marketing-and-its-influence-on-physicians-and-patients).

prescriber behavior: "[T]his prosecution gets at the heart of the matter. Patients and doctors were not, on average, irresponsible. They acted under the influence of a concerted plan of misinformation and over-promotion orchestrated up and down the supply chain for these medications."²⁷

- 74. Defendants, including Professor Pustilnik's former employer, were members of the concert and enterprise that devised and executed the plan Pustilnik identified as a primary source of the opioid crisis.
- 75. Although the introduction of OxyContin by Purdue Pharma in the late 1990's is widely acknowledged as a precipitating cause of the opioid crisis, Purdue was not the only pharmaceutical company to enthusiastically foment and exploit the booming market of controlled substances used for the treatment of pain. An industry-wide sales and marketing effort was deployed over the years by numerous manufacturers of opioid medications in order to maximize the amount of opioids they could sell.
- 76. As referenced above, OxyContin, the principal product of the Sackler family's Purdue Pharma L.P., was introduced to the market in 1996. Within six years of its introduction, the increasingly widespread misuse and abuse of OxyContin and similar opioids had drawn the attention of the United States Senate.
- 77. Two decades ago, Dr. Art Van Zee traveled from the rural coal town of St. Charles, in the southwestern corner of Virginia, to Washington D.C. to provide testimony to the United States Senate Committee on Health, Education, Labor and Pensions. On February 12, 2002, that Committee held a hearing entitled "Examining the Effects of the Painkiller OxyContin, Focusing on Federal, State, and Local Efforts to Decrease Abuse and Misuse of this Product While Assuring Availability for Patients Who Suffer Daily from Chronic Moderate to Severe Pain."²⁸

²⁷ Thomas F. Harrison, "Novel Opioid Lawsuit Goes After Ad Agency," Courthouse News, May 6, 2021, *available at:* https://www.courthousenews.com/novel-opioid-lawsuit-goes-after-ad-agency/.

²⁸ A transcript of the hearing is *available at*: https://www.govinfo.gov/content/pkg/CHRG-107shrg77770/html/CHRG-107shrg77770/html/CHRG-107shrg77770.htm

78. In those early days of the unfolding opioid epidemic, Dr. Van Zee's medical practice in St. Charles put him in a position to offer informed, first-hand observations of the toll that the pharmaceutical industry's efforts to market opioids was exacting from his community. He testified:

In the 25 years I have practiced as a general internist in St. Charles, which is a small Appalachian coal mining town, there has never been anything to compare to the epidemic of drug abuse and addiction that we have seen the last 3 years with OxyContin. Contrary to what is sometimes portrayed in the media as long-term addicts switching to the drug *du jour*, what we have seen for the most part is numerous young people recreationally using OxyContin and then becoming very rapidly addicted. Many of these kids are good kids, good families with bright, promising futures that are being destroyed in every way by their opioid addiction.²⁹

79. Further, Dr. Van Zee identified the sales and marketing practices of the pharmaceutical industry when selling controlled substances as a primary cause of the problem:

My own personal view of the complicated OxyContin abuse problem is that there are at least three major elements involved. First, there has been an obvious problem with physician misprescribing and overprescribing of this drug. Second, this epidemic has been a vicious indicator of the alarming degree of prescription drug abuse in our society. Third and perhaps the one closest to this committee and the FDA is that the promotion and marketing of OxyContin by Purdue Pharma has played a major role in this problem.³⁰

80. Five years after Dr. Van Zee's testimony and 80 miles from his hometown of St. Charles, United States Attorney John Brownlee announced in Abingdon, Virginia, the guilty plea of the Purdue Frederick Company, the parent of Purdue Pharma, L.P., relating to the misbranding of OxyContin. Brownlee stated, "Even in the face of warnings from health care professionals, the media, and members of its own sales force that OxyContin was being widely abused and causing harm to our citizens, Purdue, under the leadership of its top executives, continued to push a fraudulent marketing campaign that promoted OxyContin as less addictive, less subject to abuse, and less likely to cause withdrawal. In the process, scores died as a result of OxyContin abuse and an even greater number of people became addicted to OxyContin; a drug that Purdue led many to

³⁰ *Id.* (emphasis added).

 $^{^{29}\} See\ https://www.govinfo.gov/content/pkg/CHRG-107shrg77770/html/CHRG-107shrg77770.html/CHRG-107shrg7770.html/CHRG-107shrg7770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg770.html/CHRG-107shrg$

believe was safer, less subject to abuse, and less addictive than other pain medications on the market."³¹

- 81. Along with the guilty plea, Purdue agreed to a Corporate Integrity Agreement with the Office of Inspector General of the United States Department of Health and Human Services. For a period of five years, ending in 2012, Purdue was obligated to retain an Independent Monitor and submit annual compliance reports regarding its marketing and sales practices and training of sales representatives vis-à-vis their interactions with health care providers.
- 82. Two years later, in 2009, Dr. Van Zee published *The Promotion and Marketing of OxyContin: Commercial Triumph, Public Health Tragedy* in the American Journal of Public Health. As the title suggests, the paper applied formal rigor to some of the personal observations Dr. Van Zee previously provided to the US Senate in 2002.
- 83. In his 2009 paper, Dr. Van Zee stated the matter plainly: "Compared with noncontrolled drugs, controlled drugs, with their potential for abuse and diversion, pose different public health risks when they are overpromoted and highly prescribed." In one sense, Dr. Van Zee's observation is not particularly novel. Indeed, it approaches tautology: controlled substances are *controlled* precisely because they should not be sold to maximize volume and profits. This did not prevent Purdue and ZS from marketing its opioids full hilt, however. By 2004, "OxyContin had become the most prevalent prescription opioid in the United States."
- 84. Dr. Van Zee identified the three principal marketing tactics Purdue employed as a source of OxyContin misuse and abuse and suggested that regulation may be appropriate to curtail

³¹ See the May 10, 2007, News Release from United States Attorney, John Brownlee at https://media.defense.gov/2007/May/10/2001711223/-1/-1/1/purdue frederick 1.pdf

³² Art Van Zee, *The Promotion and Marketing of OxyContin: Commercial Triumph, Public Health Tragedy*, American Journal of Public Health, February 2009, *available at* https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2622774/pdf/221.pdf

its use. The first was the use of granular sales and marketing data to profile individual prescribers to identify those that already prescribe large amounts of opioids. "Through these profiles, a drug company can identify the highest and lowest prescribers of particular drugs in a single zip code, county, state, or the entire country. One of the critical foundations of Purdue's marketing plan for OxyContin was to target the physicians who were the highest prescribers for opioids across the country."³⁴

- 85. The second tactic was the use of incentive compensation structures to encourage the salesforce to sell ever more prescriptions of OxyContin. Bonuses at Purdue were "uncapped," meaning there was no upper limit to what an OxyContin salesperson could earn. Rather, salesforce remuneration was a direct function of overall OxyContin sales the more you sell, the more you make. "A lucrative bonus system encouraged sales representatives to increase sales of OxyContin in their territories, resulting in large numbers of visits to physicians with high rates of opioid prescriptions, as well as a multifaceted information campaign aimed at them."³⁵
- 86. The third tactic was to increase the overall number of individual calls that the salesforce placed to prescribers. "From 1996 to 2000, Purdue increased its internal sales force from 318 sales representatives to 671, and its total physician call list from approximately 33,400 to 44,500 to approximately 70,500 to 94,000 physicians." 36
- 87. When combined, these tactics produced the intended result. "The use of prescriber profiling data to target high-opioid prescribers coupled with very lucrative incentives for sales representatives would seem to fuel increased prescribing by some physicians perhaps the most liberal prescribers of opioids and, in some cases, the least discriminate." ³⁷

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- 88. Dr. Van Zee's 2002 and 2009 observations regarding the direct link between OxyContin marketing and overall opioid overdose mortality would, in time, be confirmed by further academic work, including empirical research published by the National Bureau of Economic Research in 2019.
- 89. Moreover, the legal regime under which the opioid drugs Publicis assiduously marketed and sold are regulated is the *Controlled* Substances Act. Publicis sought to maximize the sales of drugs known to be dangerous and addictive, and whose manufacture and distribution accordingly require *control*, not the same marketing tactics otherwise used for non-addictive products whose abuse liability does not routinely cause the user's death.
- 90. In 1970, Congress enacted the Controlled Substances Act ("CSA") in order to combat the spread and use of drugs known to be dangerous and/or addictive. It is also the legal regime that regulates the lawful production, possession, and distribution of substances deemed deserving of control, but that have some recognized medical use.
- 91. The Drug Enforcement Administration ("DEA") administers the act. The CSA allocates substances meriting control to one of five classifications based on the characteristics of each substance and the attendant risks posed.
- 92. In order to produce and market certain substances meriting control, pharmaceutical companies must register with the DEA and be bound by the reporting requirements of the CSA. The act requires any person who seeks to manufacture, distribute, dispense, or conduct research involving any controlled substance to obtain and maintain a registration from the DEA. *See* 21 U.S.C. 823(e); 21 C.F.R. 1301.74(b).
- 93. The opioids that Defendants and their clients marketed are classified as Schedule II controlled substances under the CSA. Schedule II substances "have a high potential for abuse for

which may lead to severe psychological or physical dependence."³⁸ As such, opioids are subject to control under the CSA because the diversion of these substances poses recognized risks to public health and safety.

- 94. The CSA also imposes reporting requirements on manufacturers, whereby registrants must monitor and report suspicious orders of opioids. These obligations include recordkeeping, whereby registrants must maintain complete and accurate inventories and records of all transactions involving controlled substances and make those records available to the DEA. In addition, registrants must periodically report all sales, delivery, disposal, or dispensing activities of any controlled substance. Schedule II controlled substance manufacturers, such as Publicis' clients, must also file Automated Reports and Consolidated Orders System (ARCOS) reports with the DEA.
- 95. Upon information and belief, Defendants, through their efforts to increase opioid sales for its clients, possessed and shared with its clients detailed information on the prescribing and dispensing patterns and volumes of its clients' Schedule II opioids. Upon information and belief, McKinsey, Publicis and ZS could determine when the prescribing or dispensing of a given clients' opioid product was unusual. For instance, upon information and belief, McKinsey, Publicis and ZS clients could identify aberrations in the number of units sold, doses prescribed, prescriptions written per prescriber, method of payment used, and other factors relevant to changes in the volume of its clients' opioid sales.
- 96. The inputs necessary for a Registrant to identify suspicious orders that merit reporting are the same inputs that McKinsey, Publicis and ZS collects, analyzes, and synthesizes in order to define and target the marketing campaigns for its clients. Upon information and belief, the

³⁸ See https://www.deadiversion.usdoj.gov/schedules/#define Schedule I substances also have a high potential for abuse and dependance. The difference is that Schedule I substances have no recognized medical use. Schedule II substances, like opioids, do.

same information utilized and analyzed by McKinsey, Publicis and ZS presented to their clients for purposes of devising and optimizing opioid sales and marketing efforts should have led to obligations by McKinsey's, Publicis' and ZS' clients – as registrants under the CSA - to report suspicious activity to the DEA.

- 97. Many states, including Oklahoma, enacted similar state laws, rules and regulations in order to regulate the manufacture, marketing, distribution and dispensing of controlled substances and provide oversight over this unique industry.³⁹
- 98. In order to keep these dangerous and addictive drugs out of the wrong hands, this closed-system of state and federal authority imposes specific duties upon Registrants to monitor, identify, halt and, perhaps most importantly, report suspicious orders of controlled substances. 21 C.F.R. § 1301.74; *Masters Pharm., Inc. v. Drug Enf't Admin.*, 861 F.3d 206 (D.C. Cir. 2017).

c. What McKinsey Does: "Consulting is more than giving advice."

- 99. McKinsey is a global consulting firm with many areas of expertise, including the pharmaceutical industry. As a management consulting firm, McKinsey provides plans to managers, directors, and owners on how to run their companies or other enterprises, and helps implement those plans.
- 100. Management consulting is the business of providing solutions to clients. Solutions take many forms, depending on the client's needs. "Management consulting includes a broad range of activities, and the many firms and their members often define these practices quite differently." 40
- 101. Broadly speaking, there are two schools of management consulting. "Strategy" consulting provides big-picture advice to clients about how they approach their business: how the business is structured, which markets to compete in, potential new business lines, and mergers and

³⁹ See 63 OK Stat. 63-2-101 et. seq.

⁴⁰ Arthur Turner, *Consulting is More Than Giving Advice*, Harvard Business Review, September 1982, *available at*: https://hbr.org/1982/09/consulting-is-more-than-giving-advice

acquisitions. The strategy consultant provides a plan to the client that the client may choose to adopt or not.

102. "Implementation" consulting is what comes next. If strategy consulting is providing advice to a client, "implementation" work is what happens once the client has adopted the consultant's plan. After a client has adopted the strategy consultant's recommendations, the implementation consultant remains in place with the client to actually do the necessary work and execute on the plan.

103. In his 1982 *Harvard Business Review* article entitled "Consulting is More Than Giving Advice," Professor Arthur Turner of the Harvard Business School described the thencurrent state of the consulting industry's attitude toward implementation work:

The consultant's proper role in implementation is a matter of considerable debate in the profession. Some argue that one who helps put recommendations into effect takes on the role of manager and thus exceeds consulting's legitimate bounds. Others believe that those who regard implementation solely as the client's responsibility lack a professional attitude, since recommendations that are not implemented (or implemented badly) are a waste of money and time. And just as the client may participate in diagnosis without diminishing the value of the consultant's role, so there are many ways in which the consultant may assist in implementation without usurping the manager's job.⁴¹

104. Although McKinsey has historically been regarded as a "strategy" consulting firm, by the time it was working with Purdue, implementation services were a core component of the suite of services that McKinsey provided within the "transformational relationship" it developed with its clients. 42 Indeed, writing in 2013, Harvard Business School Professor Clayton Christensen observed the decline in "pure" strategy work performed by consultants, as the industry sought to diversify its income streams by offering implementation and other services to clients. "For example,

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⁴² For McKinsey's own description of its implementation services, *See https://www.mckinsey.com/business-functions/mckinsey-accelerate/how-we-help-clients/implementation* (last accessed October 19, 2020).

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27 ⁴⁵ *Id.* at 308.

at traditional strategy-consulting firms, the share of work that is classic strategy has been steadily decreasing and is now about 20%, down from 60% of 70% some 30 years ago."43

105. When partnering with clients, a core component of the McKinsey relationship is discretion. "The basis of any client relationship with the firm is trust. Companies share their most competitive secrets with McKinsey with the understanding that confidentiality is paramount. McKinsey consultants aren't even supposed to tell their own spouses about their client work."44 McKinsey recognizes it must have its clients' trust and make confidentiality "paramount," as "[c]ompanies share their most competitive secrets with McKinsey" for McKinsey to do its work.⁴⁵

106. During the implementation phase, McKinsey essentially bonds with the client. Describing McKinsey's approach to implementation, one McKinsey consultant stated, "In some of the most successful engagements I've seen, you can't even tell the difference between a McKinsey team member and one of our clients because we working that cohesively together."⁴⁶

107. Another McKinsey Senior Implementation Coach described McKinsey's approach: "We're in there interacting with every element of that organization, from the welders or mechanics on the front line, all the way up to the board of directors."⁴⁷

⁴³ Clayton Christensen, Dina Wang, and Derek van Bever, "Consulting on the Cusp of Disruption," Harvard Business Review, October 2013, available at https://hbr.org/2013/10/consulting-on-the-cusp-of-disruption

⁴⁴ McDonald, *The Firm*, Pg. 308.

⁴⁶ McKinsey on Implementation, April 30, 2017, available at https://www.youtube.com/watch?v=rEQOGVpl9CY

5,507 views · Oct 22, 2018

⁴⁸ See "McKinsey Careers: what's behind McKinsey Implementation's logo and success?", October 22, 2018, available at https://web.archive.org/web/20200419140214/https://www.youtube.com/watch?v=3-Zx859VJtw ⁴⁹ Id.

108. McKinsey's implementation team even has a logo: a rowing team.



109. Jenny, a Practice Manager at McKinsey, explained its significance: "The rowers symbolized to us being in the boat with the clients, doing real work and being jointly responsible for the success." 48

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110. Eugene, a partner, explained its value:

The reason McKinsey implementation works is because clients love it. The fact that we are staying longer with them, the fact that we're getting in to the trenches, the fact that we are there to walk the emotional journey with them when they're going through the tough times and really changing their companies, is what makes McKinsey implementation truly distinctive.⁴⁹

111. In the broadest of generalities, then, McKinsey's business model, as a provider of strategy and implementation consulting services, is to partner with clients to pursue business objectives identified by McKinsey. Once an objective is identified, the client and McKinsey then

engage in concerted action as a seamless and cohesive unit in order to implement the necessary means to achieve it.

i. McKinsey's Long-Term Partnership with the Pharmaceuticals Industry

- 112. Today, McKinsey's website explains "How We Help Clients" in the pharmaceuticals industry: "Helping clients maximize commercial value by assisting with product launch, marketing, sales, and market access." McKinsey helps numerous clients throughout the pharmaceutical industry, from manufacturers to distributors and pharmacies. It often does so contemporaneously. For instance, McKinsey might advise multiple opioid manufacturers on the sales and marketing of competing branded opioid products.
- 113. McKinsey's dominance of the consulting space in the pharmaceutical industry presents its own opportunity for further client service. Specifically, McKinsey also helps its clients by telling them what their competitors—who are also McKinsey clients—are doing.
- 114. For example, McKinsey pitched its services to Purdue on the basis that it was able to "bring examples from other successful companies" and perform "detailed analytics." ⁵¹

ii. The McKinsey Pharmaceuticals and Medical Products Practice Group

- 115. Like most management consulting companies, McKinsey organizes itself into practice groups that specialize in a given industry.
- 116. McKinsey has long maintained a Pharmaceuticals and Medical Products ("PMP") industry practice group dedicated to working with pharmaceutical companies. In 2004, when McKinsey's relationship with Purdue began, the PMP group was led by Michael Pearson. Pearson worked for McKinsey for twenty-three years and was a member of the firm's shareholder council

 $^{^{50}\} https://www.mckinsey.com/industries/life-sciences/how-we-help-clients/commercial$

⁵¹ PPLPC021000601208 (emphasis added).

(McKinsey's equivalent of a board of directors) in addition to leading PMP before departing McKinsey in 2008 to helm Valeant Pharmaceuticals.⁵²

- 117. Pearson stated: "At McKinsey pharmaceuticals was one of our biggest industry groups." Pearson was "not the quintessential suave and intellectual McKinsey partner. He was loud and profane and was seen, in the words of one former colleague, as 'sharp-edged and sharp elbowed." 54
- 118. Under his leadership, McKinsey's knowledge and expertise in the pharmaceutical industry was significant. By 2009, McKinsey described its capabilities: "We have an unparalleled depth of both functional and industry expertise as well as breadth of geographical reach. Our scale, scope, and knowledge allow us to address problems that no one else can. At heart, we are a network of people who are passionate about taking on immense challenges that matter to leading organizations, and often, to the world."
- 119. In 2012, while advising Purdue, McKinsey described PMP and its health care capabilities thusly: "Indeed, there is a doctor in the house. We have more than 1,700 consultants with significant healthcare experience, including more than 150 physicians and 250 consultants with advanced degrees in genetics, immunology, biochemical engineering, neurobiology, and other life sciences. We also have 75 consultants with advanced degrees in public health, healthcare management, and related fields."

⁵² John Gapper, *McKinsey's fingerprints are all over Valeant*, Financial Times, March 23, 2016, *available at:* https://www.ft.com/content/0bb37fd2-ef63-11e5-aff5-19b4e253664a

Notably, Rob Rosiello, a McKinsey partner who was a Director of Client Services (or "DCS") of the Purdue account alongside co-DCS'es Maria Gordian and Martin Elling , went on to join Pearson at Valeant in 2015 as Chief Financial Officer. The DCS is the partner in charge of the client account.

⁵³ Michael Peltz, *Mike Pearson's New Prescription for the Pharmaceuticals Industry*, Institutional Investor, September 3, 2014, *available at:* https://www.institutionalinvestor.com/article/b14zbjfm8nf1c4/mike-pearsons-new-prescription-for-the-pharmaceuticals-industry

⁵⁴ John Gapper, *McKinsey's fingerprints are all over Valeant*, Financial Times, March 23, 2016, *available at:* https://www.ft.com/content/0bb37fd2-ef63-11e5-aff5-19b4e253664a

120. That same year, the PMP group published a report entitled "Death of a Sales Model, or Not: Perspectives on the Evolution of Pharmaceutical Field Based Selling." In it, McKinsey partner Laura Moran co-authored a segment called "The Few, The Proud, The Super-Productive: How a 'smart field force' can better drive sales." In the segment, Moran and her co-authors described various ways a pharmaceutical company could optimize its sales force. Moran worked on the Purdue account, where the strategies outlined in her article were incorporated into Project Turbocharge two years later.

- 121. With respect to pharmaceutical marketing, the PMP group states, "We support clients in creating high-impact strategies that maximize value, using customized tools. We also have detailed market data for all major geographic regions." PMP also works with pharma clients regarding their sales force: "Our efforts span the entire organization—we can help train and restructure sales forces, work directly in the field to provide coaching, maximize value from back-office services, develop strategies to accelerate short-term sales, and assist with company-wide commercial transformations." ⁵⁷
- 122. McKinsey has long considered itself a "leadership factory" for good reason.⁵⁸ Nowhere is this more apparent than the pharmaceutical industry, where, thanks to PMP's efforts under Pearson's leadership, McKinsey continues to reign as the dominant management consultant.
- 123. Consistent with PMP's ambition that McKinsey be the dominant consultant in the pharmaceutical industry, McKinsey has blanketed the entire pharmaceutical supply chain with alumni.

²⁵ Death of a Sales Model, or Not," Pharmaceutical and Medical Product Practice, McKinsey, available at https://www.mckinsey.com/~/media/mckinsey/dotcom/client_service/pharma%20and%20medical%20products/pmp%20new/pdfs/2012%20death%20of%20a%20sales%20model%20or%20not.pdf

⁵⁶ *See* https://www.mckinsey.com/industries/pharmaceuticals-and-medical-products/how-we-help-clients/commercial ⁵⁷ *Id*.

⁵⁸ See Adam Jones, "Should business schools fear McKinsey's leadership factory?," *Financial Times*, May 22, 2016, available at: https://www.ft.com/content/0d17f670-1612-11e6-b197-a4af20d5575e

124. Rajiv de Silva, for instance, was appointed CEO of Endo Pharmaceutical in March 2013. Endo's two top-selling drugs were pain medications. Endo—and de Silva, individually—have been named in multiple lawsuits related to the ongoing opioid crisis. Previously, de Silva worked with Pearson in a leadership position within PMP at McKinsey before joining Rob Rosiello, a former McKinsey partner, and Pearson at Valeant. McKinsey advised Endo on its opioid business.

- 125. Likewise, Frank Scholz was a partner at McKinsey and a leader in the PMP group for seventeen years prior to departing in 2013 to join Mallinckrodt, another opioid manufacturer presently in bankruptcy after being named in numerous lawsuits relating to the ongoing opioid crisis. In fact, Scholz was the President of the "Specialty Generics" division of Mallinckrodt (formerly SpecGX LLC), which is the division that sold generic opioids. McKinsey advised Mallinckrodt on its opioid business.
- 126. Teva Pharmaceuticals,⁶⁰ another opioid manufacturer named in numerous lawsuits for its role in the opioid crisis, is led by President and Chief Executive Officer and McKinsey alumnus, Kare Schultz. He joined the company in 2017, at which point he was also appointed to Teva's board of directors. Through an asset manager named Deerfield, McKinsey's in-house hedge fund held a financial stake in Teva Pharmaceuticals while McKinsey advised its numerous clients on how to maximize opioid sales.⁶¹

⁵⁹ David Sell, "Endo CEO downplays Valeant link," Philadelphia Inquirer, November 5, 2015, available at https://www.inquirer.com/philly/business/20151106_Endo_CEO_downplays_Valeant_link.html

⁶⁰ "Teva Pharmaceuticals" or "Teva" refers to Teva Pharmaceutical Industries Ltd. and Teva Pharmaceuticals USA, Inc., together.

⁶¹ Gretchen Morgenson, "Consulting giant McKinsey allegedly fed the opioid crisis. Now an affiliate may profit from treating addicts.," *NBC News*, February 8, 2021, *available at* https://www.nbcnews.com/news/us-news/consulting-giant-mckinsey-allegedly-fed-opioid-crisis-now-affiliate-may-n1256969 McKinsey's in-house hedge fund is discussed further, below.

127. McKinsey's involvement with Teva has been long-term. In 2006, upon his retirement from McKinsey, Roger Abravanel joined Teva's board of directors the following year.⁶² By 2011, Teva had acquired Cephalon, Inc., another manufacturer of opioids, as "a core part of [Teva's] strategy" of "growth through acquisitions."⁶³ Befitting the pattern, Cephalon had its own long-standing ties to McKinsey before being acquired by Teva. In 2008, when Cephalon's Executive Vice President, General Counsel, and Secretary John E. Osborn retired, he accepted a job as "an advisor on life sciences regulatory and compliance matters to the international consulting firm McKinsey & Company, Inc."⁶⁴

128. McKinsey has ties to another notable opioid industry combination: the 2012 acquisition of Actavis, Inc.by Watson Pharmaceuticals, Inc. ("Watson") for €4.25 billion. In the aftermath of the acquisition of the large European pharmaceutical company, Watson created a "Global Integration Management Office" reporting directly to its CEO, Paul Bisaro, to focus "on planning and implementing the integration of Actavis." In order to achieve this critical task, Watson hired Marc Lehnen: "We were very pleased to recruit Marc from McKinsey & Company, Inc. to lead the Integration Management Office. Marc has years of experience in the generic industry and knows our culture and way of operating." Notably, the press release indicates that McKinsey was already advising Watson regarding the acquisition: "Although Marc does not formally join our Company until July, he will nevertheless be involved in the integration planning during this interim period."

⁶² Form 20-F dated December 31, 2012, Teva Pharmaceutical Industries Limited, available at https://www.sec.gov/Archives/edgar/data/818686/000119312513050510/d450498d20f.htm

 $^{^{64}}$ "Cephalon General Counsel John E. Osborn to Resign Position," February 8, 2008, available at https://www.sec.gov/Archives/edgar/data/873364/000110465908008569/a08-5085_1ex99d1.htm

⁶⁵ "Watson Announces Formation of Global Integration Management Office to Support ending Actavis Acquisition," PR Newswire, May 9, 2012, *available at* https://www.prnewswire.com/news-releases/watson-announces-formation-of-global-integration-management-office-to-support-pending-actavis-acquisition-150755565.html ⁶⁶ *Id.*

⁶⁷ *Id.* (emphasis added).

129. Allergan,⁶⁸ another opioid manufacturer and defendant in the nationwide opioid litigation, has also relied on McKinsey as a source of management candidates. McKinsey Senior Adviser Christopher J. Coughlin joined Allergan's board in 2014 and remains there today.

- 130. Abbott Labs, which partnered with Purdue in the early years of OxyContin to use Abbott's sales force to market Purdue's drug, has been led by CEO Miles White since 1998. White began his career at McKinsey around 1980.
- 131. As the preceding paragraphs make clear, McKinsey was in a truly unique position: given its dominance of pharmaceutical management consulting through PMP, practically all opioid industry participants were its clients. And those same clients routinely hire McKinsey consultants to leadership positions within their companies. While advising multiple industry participants regarding the sales of competing products, McKinsey was in a position to know confidential information and trade secrets of these clients "with the understanding that confidentiality is paramount."
- 132. Because of its client relationships, McKinsey was, quite literally, the sole repository on Earth of this collective knowledge of industry-wide tactics regarding the sales and marketing of opioids, and the outcomes thereof. This unique collection of knowledge and expertise made McKinsey a hub: even if any two given industry participants did not know what each other was doing, McKinsey knew exactly what *both* were doing because both were clients.
- 133. McKinsey's relationships and influence carry far beyond the manufacturers. For instance, current McKinsey director Nancy Killefer has also been an independent director of Cardinal Health, Inc. ("Cardinal") —one of the "Big Three" Distributor Defendants in the ongoing nationwide opioid litigation—since 2015. Chunhui Moi, Cardinal's current Vice President of Corporate Strategy, was previously an associate principal at McKinsey, where he worked for nine

⁶⁸ Allergan is part of the same corporate family as Actavis and Watson.

⁶⁹ McDonald, *The Firm*, Pg. 308.

years. Michele Holcomb, Cardinal's current Executive Vice President, Chief Strategy and Business Development Officer, was a partner in the Global Pharmaceutical Practice at McKinsey.

- 134. McKinsey populates the "strategy" positions at the other opioid distributors as well. At AmerisourceBergen, the "Director of Corporate Development and Strategy" was hired away from McKinsey, where she had previously been a senior associate. AmerisourceBergen's Executive Vice President and Chief Strategy Officer had previously been a partner at McKinsey.
- 135. At McKesson Corporation ("McKesson"), another McKinsey client, the President of McKesson Specialty Health and, previously Vice President of Corporate Strategy, was Marc Owen. "Prior to joining McKesson, Owen was a senior partner at McKinsey, advising pharmaceutical manufacturers, healthcare providers, distributors and technology companies, *including McKesson*, for more than a decade." After Owen was promoted in 2012, McKesson hired yet another Vice President of Corporate Strategy away from McKinsey.
- 136. In short, one way McKinsey adds value for a client is by knowing what all of its competitors are doing. It possesses a greater body of knowledge about any given industry in which it advises multiple participants than any individual participant does itself.

iii. The Transformational Relationship

137. McKinsey has long touted the notion of a "transformational relationship." It is the goal of every client relationship McKinsey develops and, McKinsey argues, the best way to extract value from a client's use of McKinsey's services. McKinsey is not a one-off seller of advice for any given CEO's problem of the day. Rather, McKinsey argues that real value for the client derives from an ongoing "transformational" relationship with the firm.⁷¹

⁷⁰ "Marc Owen Appointed President of McKesson Specialty Health," McKesson, January 31, 2012, *available at* https://www.mckesson.com/about-mckesson/newsroom/press-releases/2012/marc-owen-appointed-president-of-mckesson-specialty-health/ (emphasis added)

⁷¹ Duff McDonald, *The Firm*, Pg. 136-37 ("McKinsey no longer pitched itself as a project-to-project firm; from this point forth [the late 1970s], it sold itself to clients as an ongoing prodder of change, the kind a smart CEO would keep around indefinitely.").

138. At its core, the "transformational relationship" is *long-term*. It is the antithesis of a one-off contract wherein McKinsey performs one discreet project for a client and then concludes its business. Rather, "once McKinsey is inside a client, its consultants are adept at artfully creating a feedback loop through their work that purports to ease executive anxiety but actually creates more of it."⁷² The long-term result can be "dependence" on the McKinsey consultants. "We insinuate ourselves," Ron Daniel, McKinsey's then-managing partner, told *Forbes* in 1987.⁷³

- 139. "They have follow-on work not just because they're good at what they do, but because they are trained in how to manage these kinds of client relationships. They understand that the core reality is the relationship and the conversation, and that any particular engagement is merely epiphenomenal," explained Alan Kantrow, formerly the editor of *McKinsey Quarterly*.⁷⁴
- 140. This strategy of weaving itself into all aspects of its clients' business proved enormously successful for McKinsey over the years. It was a strategy McKinsey encouraged its consultants to take with clients to great effect:

The sell worked: Once ensconced in the boardrooms of the biggest corporate players in the world, McKinsey rarely left, ensuring a steady and growing flow of billings for years if not decades. In 2002, for example, *BusinessWeek* noted that at that moment, the firm had served four hundred clients for fifteen years or more.⁷⁵

141. Another aspect of the transformational relationship McKinsey develops with clients is the development and marketing of "leave-behind" products, such as software applications, that are sold to clients as tools that can be used by the business on an on-going and recurring basis, separate and apart from McKinsey's project-based consulting work. As described by Harvard

⁷² *Id.* at pg. 6. Purdue provides a fine example of this feedback loop in action. In 2008, when McKinsey was advising Purdue regarding Risk Evaluation and Mitigation Strategies ("REMS") for OxyContin required by the FDA, McKinsey partner Maria Gordian wrote to fellow partners Martin Elling and Rob Rosiello regarding progress in the "REMS work" as well as "Broader Strategy work." Regarding the latter, Gordian noted that Purdue board members Jonathan Sackler and Peter Boer "basically 'blessed' [Craig Landau] to do whatever he thinks is necessary to 'save the business.'. . . *I believe there is a good opportunity to get another project here.*" MCK-MAAG-0117875 (emphasis added). Indeed, after the REMS work was completed, McKinsey continued to work on "Broader Strategy work" for another decade.

⁷³ John Merwin, "We Don't Learn from Our Clients, We Learn from Each Other," *Forbes*, October 19, 1987.

⁷⁴ Duff McDonald, *The Firm*, Pg. 185.

⁷⁵ *Id.* at pg. 136.

Business School Professor Clayton Christensen, starting in 2007, "McKinsey & Company initiated a series of business model innovations that could reshape the way the global consulting firm engages with clients. One of the most intriguing of these is McKinsey Solutions, software and technology-based analytics and tools that can be embedded at a client, providing ongoing engagement outside the traditional project-based model."

142. McKinsey's relationship with Purdue provides an example of the deployment of these "leave-behind" products. One McKinsey Solution is a pharmaceutical sales and marketing workforce optimization tool called FieldGuide, a proprietary software application McKinsey sells to clients. "The FieldGuide tool optimizes salesforce deployment and territory design through advanced geospatial analysis that leverages both market-potential insights across device categories and advanced sales-response curve analysis." McKinsey sold it to Purdue for the purpose of optimizing Purdue's OxyContin salesforce.

d. McKinsey and Purdue: A Case Study in Transformation

- 143. Indeed, McKinsey's work with Purdue is a prime example of the transformational relationship in action. McKinsey counted Purdue as a client at least as early as 2004, three years *before* Purdue's parent and officers first pleaded guilty to misbranding OxyContin in 2007. McKinsey was actively working with Purdue to increase OxyContin sales despite that guilty plea and continued to do so throughout the time period that Purdue and its advisors were bound by the terms of the Corporate Integrity Agreement entered in to alongside the guilty plea. McKinsey's work with Purdue continued through at least 2018.
- 144. McKinsey staffed at least forty known consultants to Purdue, from senior partners all the way down through engagement managers and entry-level associates. Throughout the

⁷⁶ Clayton Christensen, Dina Wang, and Derek van Bever, "Consulting on the Cusp of Disruption," *Harvard Business Review*, October 2013, *available at* https://hbr.org/2013/10/consulting-on-the-cusp-of-disruption

⁷⁷https://www.mckinsey.com/industries/pharmaceuticals-and-medical-products/how-we-help-clients/medtech/marketing-and-sales

unfolding of the nationwide opioid crisis that only continued to worsen after the 2007 guilty plea, McKinsey remained steadfast alongside the Sacklers and Purdue every step of the way. The *mea culpas* would come only later.

145. McKinsey partner Maria Gordian, in her March 26, 2009 "EY 2009 Impact Summary" internal report to McKinsey director Olivier Hamoir and McKinsey's Personnel Committee, recounted her accomplishments that year on the Purdue account. The document is an annual self-assessment produced by McKinsey partners. In it, Gordian described the state of firm's relationship for Purdue:

With client work extending through the 3rd quarter, and several additional proposals in progress, we continue to expand the depth and breadth of our relationships at Purdue. We look forward to deepening our relationships with the Sackler family and serving them on key business development issues, and to expanding our relationship with [John] Stewart and other members of the senior management team.⁷⁸

- 146. Gordian even described herself as a counselor to Richard Sackler in the same memorandum, in addition to being a "point of contact for the Board and Sackler family."⁷⁹
- 147. The continued expansion of the depth and breadth of McKinsey's relationship with Purdue was an ever-present internal goal for McKinsey, as it was accompanied by recurring and ever-increasing client billings.
- 148. By 2014, both the breadth and depth of McKinsey's relationship with Purdue had expanded dramatically. During the 2009 to 2014 period in particular, Purdue relied extensively on McKinsey to develop and implement its sales and marketing strategy for OxyContin. But McKinsey's work for Purdue involved many other facets of Purdue's business beyond sales and marketing, including general and administrative consulting, review of product acquisition,

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⁷⁸ The Ad Hoc Group of Non-Consenting States' Statement in Support of the Official Committee of Unsecured Creditors' Motions to Compel Production of Purportedly Privileged Documents for *In Camera* Review, Doc. No. 2012, *In re Purdue Pharma, Inc.*, filed November 18, 2020, Case No. 19-23649 (S.D.N.Y.), Ex 7, Pg. 48; MCK-MAAG 0118669.

⁷⁹ *Id*.

evaluation of research and development, advising Purdue on the design of clinical studies, risk management, and interactions with regulators.

149. McKinsey's sales and marketing work for Purdue focused on creating and implementing strategies and tactics to bolster the sales of OxyContin, a Schedule II drug that is widely recognized as among the most frequently diverted and abused opioids. As Purdue faced growing scrutiny, McKinsey also helped the company protect its public image and profit from the market for illicit opioids, which McKinsey's industry-wide efforts helped to promote and maintain.

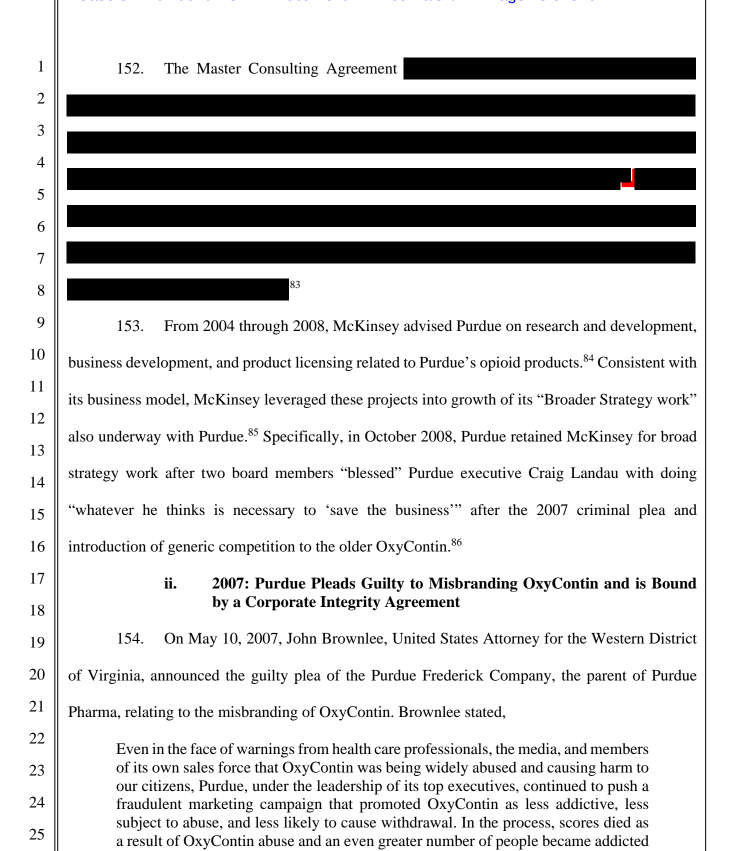
perform to maintain and grow Purdue's opioid profits amidst a growing epidemic of addiction and abuse. Part of McKinsey's work involved assessing the "underlying drivers" of OxyContin's (financial) performance. As described below, these drivers boil down to two things: (1) a widespread deceptive marketing campaign and (2) fueling an illicit market for non-medical use. Purdue entered into guilty pleas arising out of both types of conduct in 2007 and 2020, respectively. McKinsey delved into the "granular" aspects of Purdue's sales and promotion. And, throughout the two companies' long-term relationship, McKinsey understood Purdue's business "both in terms of content and culture," as its own renewed consulting agreement assured in 2013.

i. 2004: McKinsey and Purdue Meet

151. On March 1, 2004, McKinsey entered into a Master Consulting Agreement with Purdue for services that would be defined from time to time.⁸⁰ The Agreement was signed on McKinsey's behalf by Rob Rosiello, then a senior partner in the PMP practice group. After a ruling that held patents on OxyContin unenforceable due to Purdue misleading the patent office, McKinsey stepped in to help Purdue.⁸¹

⁸⁰ PPLPC012000069192

Id



⁸² *Id*.

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⁸⁴ PPLPC013000116218; PPLP004401340

⁸⁵ MCK-MAAG-0117875

⁸⁶ *Id*.

to OxyContin; a drug that Purdue led many to believe was safer, less subject to abuse, and less addictive than other pain medications on the market.

- 155. Purdue Frederick Company as well as three of Purdue's officers, pleaded guilty to the misbranding of OxyContin pursuant to various provisions of the Federal Food, Drug, and Cosmetic Act, 21 U.S.C. §§ 301, et seq.
- 156. Purdue admitted that "supervisors and employees, with the intent to defraud or mislead, marketed and promoted OxyContin as less addictive, less subject to abuse and diversion, and less likely to cause tolerance and withdrawal than other pain medications." Part of this deceptive messaging included highlighting OxyContin as a long-acting ("LA") or extended release ("ER") opioid and suggesting it created less chance for addiction than "immediate release" opioids because it had fewer "peak and trough" blood level effects or "did not cause a 'buzz' or euphoria" in the same manner as these other opioids.
- 157. Concurrent with its guilty plea, Purdue entered into a Corporate Integrity Agreement with the Office of Inspector General of the United States Department of Health and Human Services on May 7, 2007. Purdue's compliance obligations under the Corporate Integrity Agreement ran for a period of five years, and ultimately terminated in January 2013.⁸⁷
- 158. Pursuant to the Corporate Integrity Agreement, Purdue was obligated to implement written policies regarding its compliance program and compliance with federal health care program and Food and Drug Administration requirements, including:
 - a. "selling, marketing, promoting, advertising, and disseminating Materials or information about Purdue's products in compliance with all applicable FDA requirements, including requirements relating to the dissemination of information that is fair and accurate . . . including, but not limited to information concerning the withdrawal, drug tolerance, drug addiction or drug abuse of Purdue's products";
 - b. "compensation (including salaries and bonuses) for Relevant Covered Persons engaged in promoting and selling Purdue's products that are designed to ensure that financial

⁸⁷ See https://www.justice.gov/opa/press-release/file/1329576/download

incentives do not inappropriately motivate such individuals to engage in the improper promotion or sales of Purdue's products"; and

- c. "the process by which and standards according to which Purdue sales representatives provide Materials or respond to requests from [health care providers] for information about Purdue's products, including information concerning withdrawal, drug tolerance, drug addiction, or drug abuse of Purdue's products," including "the form and content of Materials disseminated by sales representatives," and "the internal review process for the Materials and information disseminated by sales representatives."
- 159. Purdue was obligated to engage an Independent Review Organization to ensure its compliance with the strictures of the Corporate Integrity Agreement and to file compliance reports on an annual basis with the Inspector General.
- 160. In the wake of its accession to the Corporate Integrity Agreement, Purdue faced newly imposed constraints on its sales and marketing practices. The Corporate Integrity Agreement was a problem to solve. Despite the agreement's constraints (i.e., do not lie about OxyContin), Purdue and its controlling owners, the Sackler family, still intended to maximize OxyContin sales.

iii. The Sacklers React to the "Concentration of Risk" Posed to Them by Purdue's Opioid Business.

- 161. The Sackler family has owned and controlled Purdue and its predecessors since 1952. At all times relevant to this Complaint, individual Sackler family members occupied either six or seven of the seats on Purdue's board of directors, and at all times held a majority of Board seats. To advise the board of directors of Purdue Pharma was to advise the Sackler family. The interests of the Sackler family and the Purdue board of directors, and Purdue itself, as a privately held company, were all aligned. Practically, they were indistinguishable.⁸⁸
- 162. As a result of the 2007 guilty plea, the Sacklers made the strategic decision to distance the family from Purdue, which was regarded, in the words of Richard Sackler, as an increasingly dangerous "concentration of risk" for Purdue's owners. Ten days after the guilty plea

⁸⁸ Craig Landau, soon to become CEO of Purdue, acknowledged in May 2017 that Purdue operated with "the Board of Directors serving as the 'de facto' CEO." The future CEO of the company, in other words, understood that he would have little practical power despite his new title. The owners ran the business.

was announced, David Sackler wrote to his father, Richard Sackler, and uncle, Jonathan Sackler, describing precisely what that "risk" was: legal liability for selling OxyContin. In response to Jonathan stating that "there is no basis to sue 'the family," David replied:

From:	David Sackler	
Sent:	5/17/2007 11:08:08 PM	
To:	'Sackier Ionathan'	I- Sackler, Dr Richard
cc:	Ives, Stephen A.	
Subject:	RE: Idea	
Attachments:	image001.jpg	

Well I hope you're right, and under logical circumstances I'd agree with you, but we're living in America. This is the land of the free and the home of the blameless. We will be sued. Read the op-ed stuff in these local papers and ask yourself how long it will take these lawyers to figure out that we might settle with them if they can freeze our assets and threaten us.

- 163. Given concern over this "concentration of risk," the two sides of the Sackler family spent considerable time and energy debating the best way to achieve distance from Purdue, and collectively considered a variety of options for doing so. One option was to sell the company to or merge the company with another pharmaceutical manufacturer. They discussed Shire as a possible target, as were Cephalon, UCB, and Sepracor, Inc. The proceeds of such a transaction could then be re-invested in diversified assets, thereby achieving the Sacklers' desired distance from opioids.
- 164. Mortimer D.A. Sackler advocated for a sale or merger in a February 21, 2008, email to Richard Sackler (a former president and co-chairman of Purdue) and several others, writing, "The pharmaceutical industry has become far too volatile and risky for a family to hold 95% of its wealth in. It simply is not prudent for us to stay in the business given the future risks we are sure to face and the impact they will have on the shareholder value of the business and hence the family's wealth." The risk he referred to was, at least in significant part, further liability related to OxyContin.
- 165. Another option was to have Purdue borrow money in order to assure Purdue had adequate funds to continue operating while the Sacklers, as owners, began to make substantial distributions of money from the company to themselves. Once again, the proceeds of the

distributions could then be re-invested in diversified assets, thereby achieving the Sacklers' desired distance.

- 166. In order to pursue either of these options, the Sacklers needed to maximize opioid sales in the short term so as to make Purdue—by then the subject of substantial public scrutiny—appear either as an attractive acquisition target or merger partner to another pharmaceutical manufacturer or as a creditworthy borrower to a lender.
- 167. In short, the Sacklers planned to engage in a final flurry of opioid pushing in order to rid themselves of their pharmaceutical company dependency for good.
- 168. In fact, in the years after the 2007 guilty plea, Purdue would retain only the absolute minimum amount of money within it as possible: \$300 million. Purdue was required to retain that amount pursuant to a partnership agreement with separate company. Otherwise, all the money was distributed to its owners.⁸⁹
- 169. Given the complexity of the problem, the Sacklers and Purdue realized that they would need assistance in achieving these internally contradictory objectives. Purdue did not have the capabilities in-house to design and implement a sales strategy for OxyContin that would achieve the Sacklers' objectives. They turned to the global management consulting firm McKinsey, which had already been advising the Sacklers and Purdue for at least three years, for help with their new problem.
- 170. Notably, under the terms of Paragraph II.C.1(b) of the Corporate Integrity Agreement, McKinsey, as a contractor to Purdue performing sales and marketing functions for the company, was itself a "Covered Person" subject to the strictures of the Agreement.⁹⁰

⁸⁹ See Jared S. Hopkins, At Purdue Pharma, Business Slumps as Opioid Lawsuits Mount, Wall Street Journal, June 30, 2019, available at: https://www.wsj.com/articles/purdue-pharma-grapples-with-internal-challenges-as-opioid-lawsuits-mount-11561887120?mod=hp_lead_pos6

⁹⁰ The relevant language in the Corporate Integrity Agreement provides: "'Covered Persons' includes . . . all contractors, subcontractors, agents, and other persons who perform sales, marketing, promotional, pricing, government contract, or regulatory functions . . . on behalf of Purdue." PDD1712900096.

9:

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iv. Purdue Tasks McKinsey with Boosting Opioid Sales in Light of the Guilty Plea and Corporate Integrity Agreement.

- 171. The Sacklers faced a problem: the need to grow OxyContin sales as dramatically as possible so as to make Purdue an attractive acquisition target or borrower, while at the same time appearing to comply with the Corporate Integrity Agreement. As one Purdue executive stated of Purdue's attitude toward the Corporate Integrity Agreement: "They did not listen to their critics and insisted they had just a few isolated problems. After the settlement, they didn't change—the way the sales force was managed and incentivized, everything stayed the same." 91
- 172. Purdue and the Sacklers were well aware of the constraints posed by the Agreement. Indeed, during a May 20, 2009 Executive Committee Meeting, the discussion led to whether Purdue should have a single sales force marketing all Purdue products, including OxyContin, or instead to "create a separate Sales Force for Intermezzo (a sleeping pill) that would be comprised of approximately 300 representatives." John Stewart, Purdue's then-CEO, saw an opportunity, and asked if the Corporate Integrity Agreement would apply if Purdue were to launch Intermezzo and another Purdue product, Ryzolt (a branded version of Tramadol, another narcotic painkiller), using the separate sales force. Might the new drug launch fall outside of the Corporate Integrity Agreement, he asked? 92
- 173. It would not, he was told by Bert Weinstein, Purdue's Vice President of Compliance. 93
- 174. Given the tension between compliance with the Corporate Integrity Agreement and the desire to sell more OxyContin, Purdue needed help.

⁹¹ David Crow, *How Purdue's 'one-two' punch fuelled the market for* opioids, Financial Times, September 9, 2018, *available at:* https://www.ft.com/content/8e64ec9c-b133-11e8-8d14-6f049d06439c

 ⁹² PPLPC012000226606, Purdue Pharma Executive Committee Meeting Notes and Actions, May 20, 2009, Pg. 2.
 93 Id

175. Ethan Rasiel, a former McKinsey consultant, has described the typical way McKinsey begins working with a client: "An organization has a problem that they cannot solve with their internal resources. That's the most classic way that McKinsey is brought in." ⁹⁴

176. Such was the case with Purdue. Because it did not have the requisite expertise to address the problems posed by the Corporate Integrity Agreement internally, Purdue expanded on its already-existing relationship with McKinsey to devise a sales and marketing strategy to increase opioid sales despite the Corporate Integrity Agreement and growing concern about the "concentration of risk" that Purdue's business of selling opioids posed to its owners.

177. McKinsey's task was to thread the needle: to increase OxyContin sales despite the strictures imposed by the five-year Corporate Integrity Agreement. This McKinsey did, turbocharging⁹⁵ the sales of a drug it knew fully well was addictive and deadly, while purporting to respect to the Corporate Integrity Agreement.

178. In short, Purdue would pay money to McKinsey in exchange for McKinsey enabling the company how to sell as much OxyContin as conceivably possible so that the Sacklers could obtain cash to diversify their investment holdings away from Purdue and keep their money safe from the reach of court judgments, fines, and penalties they feared.

179. Consistent with their plan to dissociate themselves from the company, the Sacklers appointed Mr. Stewart as the CEO of Purdue in 2007. The Sacklers viewed Stewart as someone loyal to the family. He had previously worked for a division of Purdue in Canada. Stewart's job was to assist the Sacklers with the divestiture or eventual orderly wind-down of Purdue. Stewart was paid more than \$25 million for his services to Purdue from 2007 through 2013.

⁹⁴ How McKinsey Became One of the Most Powerful Companies in the World, CNBC, June 6, 2019 available at: https://www.youtube.com/watch?v=BBmmMj_maII

⁹⁵ If the description is overbearing, note that it is McKinsey's own, as described below.

- 180. Purdue's Executive Committee discussed Stewart's concerns regarding the constraints posed by the Corporate Integrity Agreement on May 20, 2009. Within weeks, McKinsey was working with Purdue to devise and implement new marketing strategies for OxyContin.
- 181. Stewart, as CEO, was in charge of the relationship with McKinsey. He controlled workflow to and from McKinsey and required his personal approval for any work orders with McKinsey.
- 182. In addition, Purdue's Vice President of Corporate Compliance, "responsible for developing and implementing policies, procedures, and practices designed to ensure compliance with the requirements set forth in the [Corporate Integrity Agreement]," reported directly to Stewart.⁹⁶
- 183. Throughout their relationship, McKinsey routinely obtained information from, advised, communicated with, and ultimately worked for the Purdue board of directors, controlled by the Sackler family.
- 184. McKinsey would also work in granular detail with the Purdue sales and marketing staff, led during the relevant period by Russell Gasdia, Vice President of Sales and Marketing.
- 185. From as early as June 2009 and continuing at least through July 14, 2014, Purdue routinely relied upon McKinsey to orchestrate its sales and marketing strategy for OxyContin. The relationship was characterized by ongoing interactions between teams from McKinsey and Purdue regarding not only the *creation* of an OxyContin sales strategy, but also its *implementation*. McKinsey was a real presence at Purdue. "A team of McKinsey analysts went in-house, camping out in a conference room at Purdue headquarters."

⁹⁶ PDD1712900096.

⁹⁷ Patrick Radden Keefe, *Empire of Pain* 302 (2021). In September, McKinsey named Mr. Keefe's history of the Sackler family and Purdue and the opioid crisis to its 2021 shortlist for "Business Book of the Year." *See https://www.mckinsey.com/about-us/new-at-mckinsey-blog/for-your-reading-list-the-2021-business-book-of-the-year-shortlist*

98 PPLPC012000257444

⁹⁹ Ian MacDougal, *How McKinsey is Making \$100 Million (and Counting) Advising on the Government's Bumbling Coronavirus Response*, ProPublica (July 15, 2020), https://www.propublica.org/article/how-mckinsey-is-making-100-million-and-counting-advising-on-the-governments-bumbling-coronavirus-response.

100 https://www.mckinsey.com/about-us/overview

¹⁰¹ Forbes, *McKinsey* & *Company* (retrieved September 9, 2021), https://www.forbes.com/companies/mckinsey-company/?sh=1201a12624c1.

v. Purdue Relies on McKinsey.

186. Purdue hired McKinsey not only to give advice, but to devise and then implement a deceptive marketing strategy. For example, for one "major initiative" with Purdue, "McKinsey forecast[ed] a potential incremental increase in sales in the \$200-400mm range" over a three-year period, "[w]hen properly implemented." 98

187. McKinsey is not cheap, either. Indeed, hiring McKinsey is an expensive proposition. A single junior consultant—typically a recent college or business school graduate—runs clients millions of dollars annually. McKinsey is a highly selective employer and advertises that its employees join "for the opportunity to apply their talents to complex, important challenges." Talent" is key to McKinsey's model; clients pay for the best and brightest.

188. A client does not choose to pay McKinsey unless it expects to receive benefits it could not have obtained within its own organization. McKinsey offers solutions to clients facing challenges they feel they cannot adequately address on their own. This model has been a stunning success for McKinsey. In 2008, McKinsey's annual revenue was \$6 billion. Today, the firm earn more than \$10 billion in revenue each year. ¹⁰¹

189. Clients pay these exorbitant rates for a reason: McKinsey's plans and partnership work. Even critics of the consulting industry recognize the unique efficacy of McKinsey's work. JPMorgan Chase CEO Jamie Dimon once derided consultants as "substituted management" and stated that "consultants can become a disease for corporations." Dimon made one exception to this

rule: McKinsey. 102 Given unique levels of trust, respect, and access by major corporations across the United States and the world, McKinsey has unmatched power to affect how those corporations behave.

- 190. When Purdue entered into a "Master Consulting Agreement" with McKinsey in 2004, Purdue explicitly recognized McKinsey "has a fine reputation as well as excellent experience and relationships in our industry," which Purdue was counting on to boost its opioids business.¹⁰³
- 191. Purdue explicitly recognized that McKinsey stepped in to help Purdue "protect [its] sales and continue to *grow our business*." ¹⁰⁴
- 192. Furthermore, that the Sacklers, as board members of Purdue, relied on McKinsey in their conduct of Purdue affairs is an admitted fact. In a public filing in the recent Purdue bankruptcy proceedings, the one side of the Sackler family conceded that they did so: "McKinsey is widely recognized as 'a leading management consulting firm' and the Former Directors were statutorily entitled to rely on such expertise." ¹⁰⁵

vi. McKinsey Delivers.

193. Purdue, as a monoline manufacturer of opioids, relied on McKinsey in practically all aspects of its business.

1. Courting the Regulators: "We All Feel Responsible."

194. One critical aspect of Purdue's operations, given its status as a producer of controlled substances, was regulatory compliance. McKinsey guided Purdue through practically all of its interactions with regulators whose efforts to protect the public might pose threats to Purdue's business.

¹⁰² Duff McDonald. *Behind the singular mystique of McKinsey & Co.* The Guest Blog. CNBC. Sept 25, 2013. Available at: https://www.cnbc.com/2013/09/25/behind-the-singular-mystique-of-mckinsey-co.html

¹⁰³ PPLPC012000069192

¹⁰⁴ *Id.* (emphasis added).

¹⁰⁵ In re: Purdue Pharma, L.P., No. 19-23649, Doc. 3441-1, at ¶ 328 (Aug. 5, 2021).

195. McKinsey advised Purdue on how to approach the FDA in light of its criminal conviction and retain business in light of the reputational damage to the company and to OxyContin after the admissions in its guilty plea.

- 196. In 2008, Purdue submitted a New Drug Application for a reformulation of OxyContin, ostensibly to make it more difficult to abuse by extracting the active ingredient from it or otherwise defeating the time-release mechanism in OxyContin tablets—i.e., another product Purdue would later deceptively promote as safer than and less prone to abuse than it was.
- 197. Having advised Purdue on the design of tests of reformulated OxyContin as part of Purdue's FDA submission, McKinsey knew that reformulated OxyContin could still be abused. Purdue nonetheless touted its introduction of reformulated OxyContin and another ADF opioid as evidence of its good corporate citizenship and commitment to protecting the public. McKinsey worked with the Sacklers to prepare for Purdue's meetings with the FDA.
- 198. On January 20, 2009, McKinsey partner Maria Gordian wrote to partners Rob Rosiello and Martin Elling to update them on these ongoing efforts with Purdue:

We had a very good FDA rehearsal yesterday *with several family members present*. The team did an outstanding job on the study. [P]reparing the client and executing the mock meeting. We are off to DC today for the actually (sic) FDA meeting tomorrow. ¹⁰⁶

199. Gordian's email to Rosiello and Elling forwarded encouraging words from Richard Sackler. He wrote to his daughter, Marianna:

I am writing to tell you how impressed I was by the preparation for the FDA meeting. Both the method and the process as well as the content was excellent and a major departure from efforts like this in the past. Please share with the team my views and best wishes for a successful interchange with the FDA. ¹⁰⁷

Marianna forwarded the well-wishes to Gordian and the team at McKinsey.

¹⁰⁶ The Ad Hoc Group of Non-Consenting States' Statement in Support of the Official Committee of Unsecured Creditors' Motions to Compel Production of Purportedly Privileged Documents for *In Camera* Review, Doc. No. 2012, *In re Purdue Pharma*, *Inc.*, filed November 18, 2020, Case No. 19-23649 (S.D.N.Y.), Ex D, Pg. 25 (emphasis added). ¹⁰⁷ *Id.*

200. In September 2009, Purdue made a presentation to the FDA advisory committee considering its application for its reformulated OxyContin and stated that the new formulation would deter abuse. According to metadata, the PowerPoint presentation was prepared by McKinsey.

- 201. The FDA approved the reformulation of OxyContin in April 2010. 108
- 202. Having successfully navigated the approval process with McKinsey's chaperoning, Purdue then proceeded to market the ADF version of OxyContin as a solution to opioid abuse and as a reason that doctors could continue to safely prescribe their opioids.
- 203. In 2020, two FDA advisory committees evaluating the impact of the reformulated OxyContin concluded that reformulated OxyContin did not, in fact, substantially reduce abuse.
- 204. At the same time as it worked to rehabilitate Purdue's image with the FDA, McKinsey, in parallel, advised Purdue on how to limit FDA regulations aimed at mitigating the risks of opioid use. In 2008, shortly after Purdue's criminal plea, the FDA requested Purdue submit a proposed "Risk Evaluation and Mitigation Strategy" ("REMS") for OxyContin. McKinsey provided Purdue with drafts of the submission. In 109 Indeed, McKinsey was crucial in devising Purdue's response to the FDA's request for a REMS proposal from Purdue. Gordian informed Rosiello and Elling on October 23, 2008, that John Stewart, Purdue's CEO, "is aware of the critical role we are playing in pulling REMs together and is very appreciative." In the same email, she noted that "the family" was focused "on the response to the non-approval letter" from the FDA. In 110
- 205. In 2009, the FDA expanded its scope to a class-wide extended release/long-acting REMS program.

 $^{^{108} \}it See \ https://www.fda.gov/media/126835/download$

¹⁰⁹ PDD8901578031

¹¹⁰ The Ad Hoc Group of Non-Consenting States' Statement in Support of the Official Committee of Unsecured Creditors' Motions to Compel Production of Purportedly Privileged Documents for *In Camera* Review, Doc. No. 2012, *In re Purdue Pharma, Inc.*, filed November 18, 2020, Case No. 19-23649 (S.D.N.Y.), Ex C, Pg. 22.

206. Seeking to avoid a requirement that prescribers undergo mandatory training on OxyContin's risks or management or obtain certification before prescribing OxyContin, which would limit the numbers of available prescribers, Purdue turned to McKinsey. McKinsey found the cost to Purdue of a system to verify completion of prescriber education before prescriptions could be filled would be \$50 million—an estimate Purdue used to oppose efforts for more rigorous risk management strategies.¹¹¹

207. Armed with McKinsey's analysis, Purdue's strategy on REMS was effective. The

REMS program avoided verification and enrollment provisions that would harm Purdue's profits.

208. Meanwhile, based on McKinsey's work on extended-release opioid REMS, McKinsey was aware of warnings and adverse events included within the OxyContin medication guide and communications plans, including risks of overdose and adverse events including dizziness and lethargy.

209. In June 2009, McKinsey helped Purdue prepare for an FDA advisory committee meeting.

210. McKinsey prepared for Purdue an "FDA Advisory Committee on Reformulated OxyContin: Question & Answer Book" in September 2009, with questions including "Why should we trust you?" In response, McKinsey recommended Purdue say "We acknowledge mistakes made in the past[;]" "We have x, y and z measures in place that did not exist before[;]" and "[a]t all levels,"

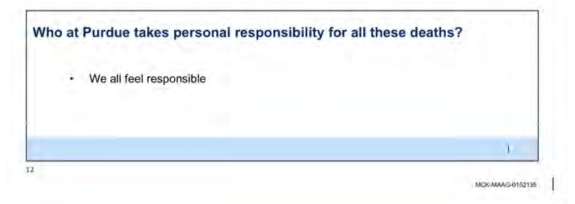
¹¹¹ PDD8901530124

¹¹² PPLPC019000622253

¹¹³ PDD8901645845

Purdue's focus is on maintaining the highest ethical standards and meeting the needs of patients[.]"114

211. Sometimes, McKinsey's work was as obfuscating as it was self-revealing. To the question of "Who at Purdue takes personal responsibility for all these deaths?[,]" McKinsey offered the following response: 115



2. The Granularity of Growth

- 212. To this end, McKinsey prides itself on certain managerial techniques it professes to have detailed knowledge of and expertise in deploying. These techniques are generally applicable to problems encountered by many businesses; they are conceptual frameworks that McKinsey deploys when tasked with solving a problem for a client.
- 213. After Purdue's first guilty plea, the Sacklers desired dramatic, short-term growth of Purdue's opioid sales so as to increase the company's attractiveness as an acquisition target or borrower while allowing the Sacklers to take money out of the company. One service McKinsey offers to its clients is to tell them how to grow.
- 214. In order to identify growth opportunities for a client, McKinsey espouses a "granular" approach to identifying which subsets of the client's existing business are the sources

¹¹⁴ MCK-MAAG-0152135

¹¹⁵ The Ad Hoc Group of Non-Consenting States' Statement in Support of the Official Committee of Unsecured Creditors' Motions to Compel Production of Purportedly Privileged Documents for *In Camera* Review, Doc. No. 2012, *In re Purdue Pharma, Inc.*, filed November 18, 2020, Case No. 19-23649 (S.D.N.Y.), Ex F, Pg. 39.

of growth, and exploiting them for all they are worth. In August 2008, McKinsey directors Patrick Viguerie and Sven Smit, together with Mehrdad Baghai, published a treatise on the matter: *The Granularity of Growth: How to Identify the Sources of Growth and Drive Enduring Company Performance* (Wiley, April 2008). "The key is to focus on granularity, to breakdown big-picture strategy into its smallest relevant components." ¹¹⁶

215. Previously, in an article in *McKinsey Quarterly* (coincidentally published the same month that Purdue pled guilty), the authors explained:

Our research on revenue growth of large companies suggests that executives should "de-average" their view of markets and develop a granular perspective on trends, future growth rates, and market structures. Insights into subindustries, segments, categories, and micromarkets are the building blocks of portfolio choice. Companies will find this approach to growth indispensable in making the right decisions about where to compete. ¹¹⁷

- 216. Additionally, McKinsey encouraged a granular assessment of the geography of corporate growth. "The story gets more precise as we disaggregate the company's performance on the three growth drivers in 12 product categories for five geographic regions." ¹¹⁸
- 217. One can imagine this strategy applied to a seller of, say, cartons of milk. If McKinsey were to perform an analysis of the milk seller's sales and marketing and discover that the profit margin on milk cartons sold to university cafeterias in dairy-producing states is much greater than the margin on cartons sold at convenience stores in the southwest, and further that the milk seller has previously devoted equal amounts of time and resources selling to both university cafeterias and convenience stores, then McKinsey would likely advise the client to deploy additional resources towards selling milk to university cafeterias in dairy-producing states.

The granularity of growth, Book Excerpt, McKinsey & Company, March 1, 2008, available at: https://www.mckinsey.com/business-functions/strategy-and-corporate-finance/our-insights/the-granularity-of-growth Mehrdad Baghai et. al., The granularity of growth, McKinsey Quarterly, May 2007, available at: https://www.mckinsey.com/featured-insights/employment-and-growth/the-granularity-of-growth

McKinsey's "granular" approach to the milk seller's business channels has identified a way to increase higher margin sales, leading to newfound growth and profitability for the client.

218. Rather than milk, McKinsey deployed this strategy on OxyContin, a controlled substance, after its manufacturer pled guilty to misrepresenting the addictive and deadly properties of the drug.

3. "Identifying Granular Growth Opportunities for OxyContin"

- 219. McKinsey's granular analysis of Purdue's OxyContin sales efforts led to the implementation of a number of strategies to sell more pills.
- 220. By January 2010, McKinsey informed Purdue, in accordance with the lessons of McKinsey's granular growth analysis, that Purdue could generate "\$200-400mm" in additional annual sales of OxyContin by implementing McKinsey's strategies.¹¹⁹
- 221. In November 2010, a McKinsey report instructed sales reps to maximize profits by "emphasizing [the] broad range of doses"—which meant pushing the doses that were highest and most profitable. 120
- 222. In 2012, John Stewart assigned McKinsey to "understand the significance of each of the major factors affecting OxyContin's sales." ¹²¹
- 223. This McKinsey did in excruciatingly granular detail, analyzing each sales channel for Purdue's opioids to identify weaknesses, opportunities, and to suggest courses of action to improve performance. Many core themes of McKinsey's work would be crystallized in a series of presentations and updates made to the Sackler family and to Purdue's board of directors in the summer of 2013 entitled "Identifying Granular Growth Opportunities for OxyContin."

a. Marketing – Countering Emotional Messages

- 56 -

¹¹⁹ PPLPC012000257443; PPLPC012000257446

¹²⁰ PPLPC018000346294

¹²¹ PPLPC020000587064

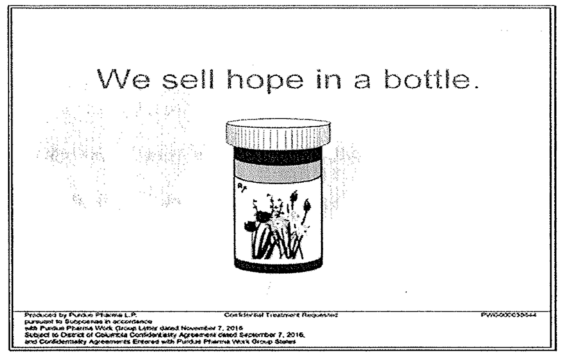
224. From the outset of McKinsey's known work for Purdue, the work was grim. In June 2009, McKinsey teamed with Purdue's then-Chief Medical Officer (and current CEO) Craig Landau and his staff to discuss how best to "counter emotional messages from mothers with teenagers that overdosed in [sic] OxyContin."

- 225. Months later, McKinsey advised Purdue to market OxyContin based on the false and misleading notion that the drug can provide "freedom" and "peace of mind" for its users, give patients "the best possible chance to live a full and active life," and concomitantly reduce stress and isolation.¹²²
- 226. These marketing claims were tailored to avoid any pitfalls that the Corporate Integrity Agreement might hold. While false and misleading, these claims regarding "freedom" and "peace of mind" of OxyContin users were narrowly tailored in order to avoid representations regarding "the withdrawal, drug tolerance, drug addiction or drug abuse of Purdue's products," as specified in Section III.B.2.c of the Corporate Integrity Agreement.¹²³

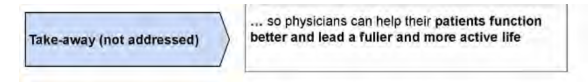
¹²² PPLPC023000239858

 $^{^{123}\} PDD1712900096$

227. Purdue's marketing materials from that time period are illustrative of the approach: 124



228. Likewise, McKinsey informed Purdue that by highlighting the ability to "tailor the dose" and treat a "broad range of appropriate patients," the prescriber-takeaway would be that "physicians can help their patients function better and lead a fuller and more active life," even though this conclusion was not to be explicitly addressed.¹²⁵



229. Claims that OxyContin improved function and quality of life were not supported by substantial evidence and, in addition, failed to take into account risks of addiction. The FDA and other federal agencies have, for years, made clear the lack of evidence for claims that the use of opioids for chronic pain improves patients' function and quality of life. A Centers for Disease

¹²⁴ Tennessee v. Purdue Pharma L.P., Case No. 1-173-18 (Compl. May 15, 2018) ¶ 24.

¹²⁵ PPLPC019000329253

¹²⁶ The FDA has warned other drug makers that claims of improved function and quality of life were misleading. See

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Control and Prevention guideline, following a "systematic review of the best available evidence," concluded that "[w]hile benefits for pain relief, function and quality of life with long-term opioid use for chronic pain are uncertain, risks associated with long-term opioid use are clearer and significant." According to the CDC director, "for the vast majority of patients, the known, serious, and too-often- fatal risks far outweigh the unproven and transient benefits [of opioids for chronic pain]." ¹²⁸

- 230. In addition to crafting carefully-tailored quality of life assurances designed to avoid the pitfalls of the Corporate Integrity Agreement, McKinsey invented other misleading marketing efforts for Purdue.
- 231. For instance, McKinsey urged Purdue to capitalize on OxyContin's extended-release characteristics in another way: marketing OxyContin's twelve-hour dosing as though users only need to take OxyContin twice a day, thus requiring fewer pills. OxyContin in fact was well known to wear off after eight to ten hours in many patients, however. What McKinsey called "convenient," would later be called "a [d]escription of Hell."
- 232. This misleading assurance of twelve-hour relief is especially pernicious, as end-of-dose failure renders OxyContin even more dangerous because patients begin to experience withdrawal symptoms, followed by a euphoric rush with their next dose—a cycle that fuels a craving for OxyContin. For this reason, Dr. Theodore Cicero, a neuropharmacologist at the

Warning Letter from Thomas Abrams, Dir., FDA Div. of Mktg., Adver., & Commc'ns, to Doug Boothe, CEO, Actavis Elizabeth LLC (Feb. 18, 2010) (rejecting claims that Actavis' opioid, Kadian, had an "overall positive impact on a physical and mental functioning, daily activities, or enjoyment of life."). work, ALLERGAN MDL 00387583; Warning Letter from Thomas Abrams, Dir., FDA Div. of Mktg., Adver., & Commc'ns, to Brian A. Markison, Chairman, President and Chief Executive Officer, King Pharmaceuticals, Inc. (March 24, 2008) (finding the claim that "patients who are treated with [Avinza (morphine sulfate ER)] experience an improvement in their overall function, social function, and ability to perform daily activities . . . has not been demonstrated by substantial evidence or substantial clinical experience."). ALLERGAN CA 00161496. The FDA's warning letters were available to McKinsey on the FDA website. ¹²⁷ CDC Guideline at 2, 18.

¹²⁸ Thomas R. Frieden and Debra Houry, New England Journal of Medicine, "Reducing the Risks of Relief – The CDC Opioid-Prescribing Guideline" at 1503 (Apr. 21, 2016).

Washington University School of Medicine in St. Louis, called OxyContin's twelve-hour dosing "the perfect recipe for addiction." Many patients will exacerbate this cycle by taking their next dose ahead of schedule or resorting to a rescue dose of another opioid, increasing the overall amount of opioids they are taking. Promotion of twelve-hour dosing, without disclosing its limitations, is misleading because it implies that the pain relief supplied by each dose lasts twelve hours.

- 233. In addition to designing misleading marketing messages, McKinsey even suggested encouraging a new channel through which those messages could be delivered to prescribers. McKinsey encouraged the tactic of "patient pushback," wherein McKinsey and Purdue would foment patients to directly lobby their doctors for OxyContin even when those physicians expressed reservations regarding the administration of Purdue's opioids.
- 234. The idea was that McKinsey and Purdue could spread their own message through pain patients who would be perceived as more credible sources suggesting a need for controlled or extended-release opioid—even though the team devising this strategy would have known that extended-release opioids did not substantially control pain or thwart addiction better than lower-dose, immediate release opioids.¹³⁰
- 235. McKinsey also coached Purdue on building "trust" (which from its vantage point, McKinsey knew was misplaced) in Purdue following its criminal conviction.

b. Targeting – Selling More OxyContin to Existing High Prescribers

236. Perhaps the key insight McKinsey provided was, using its granular approach, to identify historically large prescribers and target ever more sales and marketing resources on them, without any regard for, and indeed conscious disregard of, patient safety. Physician targeting proved effective. McKinsey advised Purdue that visiting high-prescribing doctors many times per

¹²⁹ Harriet Ryan, "''You Want a Description of Hell?' OxyContin's 12-Hour Problem," Los Angeles Times, May 5, 2016, available at http://www.latimes.com/projects/oxycontin-part1/.

¹³⁰ PDD8901645845

year increased sales. This relentless drive to increase sales and create greater availability of opioids was made with no notable concern about the parallel increase in opioid-related deaths, abuse, and misuse.

- 237. On January 20, 2010, Purdue's board was informed of the ongoing work McKinsey was performing concerning a new "physician segmentation" initiative whereby McKinsey would analyze the opioid prescribing patterns of individual physicians to identify those that had historically been the highest prescribers. McKinsey then worked with Purdue's sales and marketing staff to specifically target those prescribers with a marketing blitz to encourage even further prescribing.
- 238. Purdue trained its sales force in tactics to market to these high prescribers based on McKinsey's insights and designed in conjunction with McKinsey.
- 239. Many of the historically highest prescribers of OxyContin—those same individuals that McKinsey urged Purdue to target for ever more prescriptions—had prescribed Purdue's OxyContin *before* the 2007 guilty plea and had already been subjected to Purdue's misrepresentations regarding OxyContin that were the subject of that guilty plea.
- 240. McKinsey identified these physicians—those that had already been influenced by Purdue's misrepresentations and were thus already high prescribers—as optimal targets for a massive marketing push to sell more OxyContin.
- 241. McKinsey worked assiduously with Purdue over many years to continually refine this approach and required ever-more granular data for its analysis. More than three years after the initial introduction of the physician segmentation initiative, McKinsey requested, and Purdue provided, "prescriber-level milligram dosing data" so that McKinsey could further analyze the individual amounts of OxyContin prescribed by individual physicians.

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¹³¹ PPLPC012000257446

- 242. At the same time, it requested this "prescriber-level milligram dosing data" from Purdue, McKinsey urged the Sacklers to strictly manage the target lists of each sales representative to assure that the maximum amount of each sales representative's time was spent with the most attractive customers.
- 243. On July 23, 2013, Purdue's board discussed concerns about "the decline in higher strengths" of Purdue's opioids as well as an observed decline is "tablets per Rx." In order to assure that the threat to OxyContin sales growth be addressed, McKinsey was assigned "to actively monitor the number and size of opioid prescriptions written by individual doctors." ¹³²
- 244. In unveiling Project Turbocharge to Purdue and the Sacklers, McKinsey stated that the most prolific OxyContin prescribers wrote "25 times as many OxyContin scripts" as less prolific prescribers and urged Purdue and the Sacklers to "make a clear go-no go decision to 'Turbocharge the Sales Engine'" by devoting substantial capital toward McKinsey's plan.¹³³
- 245. McKinsey also stated that increased numbers of visits by sales representatives to these prolific prescribers would increase the number of opioid prescriptions that they would write. This singular focus on increasing prescriptions was not coupled with colorable concern for the patient population.
- 246. By November 2013, McKinsey had obtained the physician-level data they had previously requested and continued to study ways to sell additional OxyContin prescriptions by refining and targeting the sales pitch to them. The Purdue board was kept apprised of McKinsey's progress.
- 247. Not only did McKinsey identify which doctors prescribed the most of Purdue's opioids, McKinsey also recommended segmenting prescribers into "types" and tailoring messages and tactics to the different prescriber profiles. For prescribers dubbed "Early Adopting Experts"

¹³² PPLP004307354

¹³³ PPLP004409890

and "Proactive Teachers," defined by a willingness to use extended release opioids, including in opioid naïve patients (patients who were not already using opioids), McKinsey urged emphasizing that its seven tablet strengths provide flexibility to "tailor the dose" to customer needs. ¹³⁴ Upon information and belief, this message aimed to encourage prescribers to initiate and maintain patients on OxyContin long-term by reminding them they could increase the dose as patients became tolerant with long-term use (rather than discontinue use when the drug lost its effectiveness).

- 248. In its October 26, 2009, presentation, "OxyContin driving growth through stronger brand loyalty," McKinsey proposed tactics to turnaround declining sales, "[e]nhance loyalty to OxyContin among loyalist prescribers," "[c]onvert[ing] 'fence sitters' into more loyal OxyContin prescribers," and "[p]rotect OxyContin's market share[.]" In other words, McKinsey proposed increasing sales by pushing both willing and reluctant physicians to prescribe more OxyContin.
- 249. McKinsey also recommended a strategy to target those prescribers who did not regularly prescribe OxyContin—so-called "Resigned Followers and ER Delayers" —encouraging them to "increase step-up" to extended-release opioids. These were physicians with "low comfort with extended-release opioids." McKinsey encouraged Purdue to emphasize to them the "range of appropriate patients." In other words, McKinsey's strategy recommended that Purdue encourage prescribers to use OxyContin earlier in a patient's treatment for a wider range of patients and for longer periods of time.

c. Titration – Selling Higher Doses of OxyContin

250. McKinsey understood that the higher the dosage strength for any individual OxyContin prescription, the greater the profitability for Purdue. Of course, higher dosage strength,

¹³⁴ MCK-MDL2296-0126522

¹³⁵ Id

¹³⁶ *Id.* at 2.

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particularly for longer periods of use, also contributes to opioid dependency, addiction and abuse. Nonetheless, McKinsey advised Purdue to focus on selling higher strength dosages of OxyContin.

- 251. Consistent with its granular growth analysis, as early as October 26, 2009, McKinsey advised the Sacklers and the Purdue board that Purdue should train its sales representatives to "emphasiz[e] the broad range of doses," which would have the intended effect of increasing the sales of the highest (and most profitable) doses of OxyContin. ¹³⁷
- 252. McKinsey's work on increasing individual prescription dose strength continued throughout the time period McKinsey worked with Purdue. The Sacklers were informed on July 23, 2013, that Purdue had identified weakness in prescribing rates among the higher doses of OxyContin and reassured the Sacklers that "McKinsey would analyze the data down to the level of individual physicians" in order to study ways to maximize the sales of the highest-dose OxyContin pills.
- 253. Purdue implemented McKinsey's suggestions through adopting the marketing slogan to "Individualize the Dose" and by 2013 encouraged its sales representatives to "practice verbalizing the titration message" when selling Purdue's opioids to prescribers. 138
- 254. McKinsey would have known, however, that higher doses of opioids carry greater risk. Patients receiving high doses of opioids (e.g., doses greater than 100 mg morphine equivalent dose ("MED") per day) as part of long-term opioid therapy are three to nine times more likely to suffer overdose from opioid-related causes than those on low doses. As compared to available alternative pain remedies, scholars have suggested that tolerance to the respiratory depressive effects of opioids develops at a slower rate than tolerance to opioids' analgesic effects. The Centers

¹³⁸ PPLP003450924

for Disease Control and Prevention also recognize that higher doses of opioids tend to increase overdose risks relative to any potential patient benefit. 139

255. Claims that opioids could be taken in ever-increasing strengths to obtain pain relief, without disclosing that higher doses increased the risk of addiction and overdose, are deceptive and misleading. They were particularly important to promotional efforts, however, because patients on opioids for more than a brief period develop tolerance, requiring increasingly high doses to achieve pain relief. Marketers needed to generate a comfort level among doctors to ensure the doctors maintained patients on the drugs even at the high doses that became necessary.

Purdue adopted McKinsey's proposal. 140 256.

257. The titration messaging worked. Nationwide, based on an analysis by the Los Angeles Times, more than 52% of patients taking OxyContin longer than three months are on doses greater than sixty milligrams per day, which converts to the ninety MED that the CDC guideline urges prescribers to "avoid" or "carefully justify." ¹⁴²

> d. Covered Persons - Sales Quotas and Incentive Compensation

PPLPC012000246009 (PPLPC021000265092

¹³⁹ Dowell D, Haegerich TM, Chou R. CDC Guideline for Prescribing Opioids for Chronic Pain – United States, 2016. MMWR Recomm Rep 2016;65(No. RR-1):1-49. DOI: http://dx.doi.org/10.15585/mmwr.rr6501e1 see also

PPLPC023000251226 PPLPC012000245087 PPLPC012000243668 (

^{.41} PKY183123435

¹⁴² CDC Guideline at 16.

258. McKinsey urged the use of quotas and bonus payments to motivate Purdue's sales force to sell as many OxyContin prescriptions as possible. As McKinsey described it, "[r]evision to incentive comp could better align reps to Purdue's economics." 143

- 259. Notably, this behavior was prohibited by the 2007 Corporate Integrity Agreement, which required Purdue to implement written policies regarding "compensation (including salaries and bonuses) for [sales representatives] engaged in promoting and selling Purdue's products that are designed to ensure that financial incentives *do not inappropriately motivate such individuals to engage in the improper promotion or sales of Purdue's products*."¹⁴⁴
- 260. By 2010, Purdue had implemented a four-year plan, consistent with McKinsey's strategy, to dramatically increase the quota of required annual sales visits by Purdue sales representatives to prescribers. The quota was 545,000 visits in 2010, 712,000 visits in 2011, 752,000 in 2012, and 744,000 visits in 2013.
- 261. On August 15, 2013, as part of their "Identifying Granular Growth Opportunities for OxyContin" presentation, McKinsey urged the Sacklers to "establish a revenue growth goal (e.g., \$150M incremental stretch goal by July 2014) and set monthly progress reviews with CEO and Board."¹⁴⁵
- 262. In its "Identifying Granular Growth Opportunities for OxyContin" presentation to the Purdue board in July 2013, McKinsey nonetheless urged Purdue, in addition to increasing the focus of the sales force on the top prescribers, to increase the overall quotas for sales visits for individual sales representatives from 1,400 to 1,700 annually.

¹⁴³ PPLPC012000441016

¹⁴⁴ PDD1712900096 (emphasis added).

¹⁴⁵ PPLP004409890

263. In 2013, McKinsey identified one way that Purdue could squeeze more productivity out of its sales force: by slashing *one third* of the time that Purdue devoted to training its sales force (from 17.5 days per year to 11.5 days):

One possible way to attain benchmark ~1500 calls per year is to decrease training days by -6 days and increase calls per day by 5% . One possible route to benchmark Current call activity Potential new allocation Number of "on territory" days per year Number of "on territory" days per year Days1 Days1 Number of working days 260 Number of working days 260 -11.3 Holidays Holidays Vacation and other time off -27.2 Vacation and other time of -27.2 Trainings and meetings -17.5 Trainings and meetings Other company-related time off of field Other company-related time off of field -4.3 -4.3 Total days 199.7 Total days 205.7 Avg calls per day 7 Avg calls per day × 1398 Total calls per year Total calls per year 1 Purdue 2012 Actual data was used for this analysis McKessey & Company SOURCE: Purdue: team analysis

264. By eliminating one third of the amount of time sales representatives were required to be in training, McKinsey projected that Purdue could squeeze an additional 5% of physical calls per day out of its newly less-trained sales force.

265. Additionally, McKinsey developed and advised Purdue on a new incentive compensation structure for the sales representatives, who were Covered Persons pursuant to the Corporate Integrity Agreement. McKinsey knew that, combined with the strictures of sales quotas and less training for the sales force, bonus/incentive compensation to the sales representatives based on the number of OxyContin prescriptions the representative produced could be a powerful driver of incremental OxyContin sales, without regard for patient safety.

vii. Transformation: Purdue and McKinsey Adopt and Implement McKinsey's Strategies.

- 266. As early as September 11, 2009, McKinsey determined and told Purdue that it could generate \$200 million to \$400 million in additional annual sales of OxyContin by implementing McKinsey's strategy based on the opportunities its granular growth analysis had identified. McKinsey reiterated its assurances regarding the hundreds of millions of dollars of additional OxyContin sales on January 20, 2010.
- 267. Purdue accepted and, with McKinsey's ongoing assistance, implemented McKinsey's strategies for selling and marketing OxyContin.
- 268. For instance, in January 2010, Purdue was training its sales and marketing force on the new sales tactics based on a "physician segmentation" initiative that McKinsey urged. The strategy developed as a result of McKinsey's granular analysis of OxyContin sales channels. The initiative sought to identify the most prolific OxyContin prescribers and then devote significant resources towards convincing those high prescribers to continue to prescribe ever more OxyContin, in higher doses, for longer times, to ever more patients.
- 269. On January 20, 2010, the Purdue board was informed of the progress in implementing McKinsey's "physician segmentation" initiative.
- 270. This transformative collaboration would continue over the course of the relationship between Purdue and McKinsey.
- 271. During the time that McKinsey was working with Purdue, Purdue deliberately minimized the importance of the Corporate Integrity Agreement. In 2008, Carol Panara joined the Purdue sales force from rival Novartis. She would stay with the company until 2013, during which time McKinsey was responsible for increasing OxyContin sales at Purdue, and culminating with the implementation of McKinsey's "Project Turbocharge," beginning September 2013.
- 272. Ms. Panara stated that the 2007 guilty plea was deliberately minimized by the company in presentations to its sales staff: "They said, 'We were sued, they accused us of mis-

28 | 147 *Id.* 148 PPLPC030000770531

marketing, but that wasn't really the case. In order to settle it and get it behind us we paid a fine.' You had the impression they were portraying it as a bit of a witch hunt."¹⁴⁶ (Purdue and its executives paid \$634.5 million in fines.)

273. Consistent with McKinsey's mandate, McKinsey devised methods for sales staff to sell OxyContin to doctors while at the same time maintaining technical compliance with the Corporate Integrity Agreement. Ms. Panara stated that, though she was told she could not flatly claim that OxyContin was better or safer than other opioids, "she was trained to talk about product in ways that implied that it was safer." She might tout OxyContin's twelve-hour formulation to a prescriber. "You could say that with a shorter-acting medication that wears off after six hours, there was a greater chance the patient was going to jump their dosing schedule and take an extra one a little earlier. We couldn't say [it was safer], but I remember we were told that doctors are smart people, they're not stupid, they'll understand, they can read between the lines." 147

viii. Project Turbocharge a/k/a Evolve to Excellence a/k/a E2E

- 274. The Corporate Integrity Agreement expired in January 2013. With this restriction lifted, McKinsey devised additional marketing and sales strategies for Purdue to further increase OxyContin sales.
- 275. On May 14, 2013, McKinsey entered into a "Statement of Services to the Master Consulting Agreement" (the "2013 Agreement") with Purdue to "conduct a rapid assessment of the underlying drivers of current OxyContin performance, identify key opportunities to increase near-term OxyContin revenue and develop plans to capture priority opportunities."

¹⁴⁶ David Crow, *How Purdue's 'one-two' punch fuelled the market for opioids*, Financial Times, September 9, 2018, available at: https://www.ft.com/content/8e64ec9c-b133-11e8-8d14-6f049d06439c

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¹⁵¹ *Id*. 27 152 PPLPC012000431809

¹⁴⁹ *Id*.

153 PPLPC012000431262

154 Id.; PPLPC012000431266

150 PPLPC030000770531 / MCK-MAAG-0024283

276. The 2013 Agreement stated, "We have a long history of partnership with Purdue, and we would make best efforts to leverage our understanding of your business – both in terms of content and culture." It was signed by then-principal Arnab Ghatak, who would "lead the team with senior leadership from Rob Rosiello and Martin Elling." Elling was a leader of McKinsey's PMP group. 149

McKinsey was tasked with "Identifying Granular Growth Opportunities for 277. OxyContin," conducting an "assessment of the underlying drivers of current OxyContin performance," identifying "key opportunities to drive near-term OxyContin performance," and developing "plans to capture priority opportunities." ¹⁵⁰

278. For purposes of the project, McKinsey would need "[f]ull access to work done to date and key data."151 And,

279. Staff told the Sacklers that McKinsey would study how to get doctors to prescribe more OxyContin, ¹⁵³ how to use incentive compensation to push reps to generate more prescriptions, how to use "patient pushback" to get doctors to prescribe more opioids, and how to keep patients on opioids longer. 154

The 2013 Agreement would lead to Project Turbocharge, McKinsey's successful 280. bid to transform Purdue's sales and marketing efforts for OxyContin now that Purdue was no longer bound by the Corporate Integrity Agreement.

281. In the summer of 2013, McKinsey made multiple recommendations to Purdue's board to increase OxyContin revenue, and urged the Sackler family to "make a clear go-no-go decision to 'Turbocharge the Sales Engine."

- 282. Purdue, like McKinsey, recognized that the initiative was no small thing. An internal Purdue email states that
- 283. The Sacklers were impressed. On August 15, 2013, Richard Sackler emailed Mortimer D.A. Sackler, "[T]he discoveries of McKinsey are astonishing."
- 284. Eight days later, on August 23, 2013, McKinsey partners met with the Sackler family—not the Purdue board of directors—to pitch Project Turbocharge. Dr. Arnab Ghatak, one of the McKinsey partners leading the Purdue account, recounted the meeting to fellow McKinsey partner Martin Elling in an email exchange: "[T]he room was filled only with family, including the elder statesman Dr. Raymond [Sackler] We went through exhibit by exhibit for about 2 hrs They were extremely supportive of the findings and our recommendations . . . and wanted to strongly endorse getting going on our recommendations." ¹⁵⁶
- 285. Elling, a co-leader of the Purdue account, remarked in the same email correspondence that McKinsey's "findings were crystal clear to" the Sacklers, and that the Sacklers "gave a ringing endorsement of 'moving forward fast." ¹⁵⁷
- 286. As a result of the Sackler family endorsement of McKinsey's proposals, the following month Purdue implemented Project Turbocharge based on McKinsey's recommendations. In adopting "Project Turbocharge," Purdue acknowledged the improper

¹⁵⁷ *Id*.

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¹⁵⁵ PPLPC012000437344

¹⁵⁶ MCK-MDL2996-0403095

connotations of the name, and re-christened the initiative the decidedly more anodyne "E2E: Evolve to Excellence." ¹⁵⁸

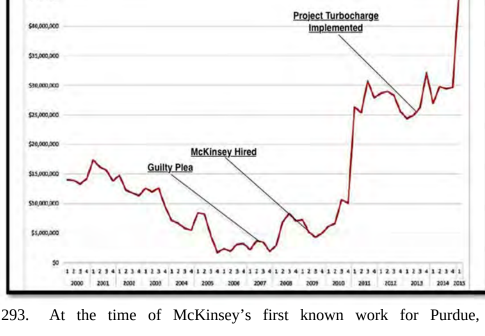
- 287. Evolve to Excellence ("E2E") was the theme of Purdue's 2014 National Sales Meeting.
- 288. CEO John Stewart also told sales staff that board member Paolo Costa was a "champion for our moving forward with a comprehensive 'turbocharge' process," referring to McKinsey's plan.
- 289. After Purdue adopted McKinsey's recommendations, McKinsey continued to work with Purdue sales and marketing staff reporting to Russell Gasdia during Purdue's implementation of McKinsey's recommendations.
- 290. In fact, the entire E2E initiative was overseen by McKinsey and some Purdue executives, who together comprised the E2E Executive Oversight Team and Project Management Office.
- 291. At the same time, the Sacklers were kept informed of the implementation of McKinsey's OxyContin strategy. According to a September 13, 2013, board agenda, the board discussed with the Sacklers the ongoing implementation of McKinsey's sales tactics.
- 292. Evolve to Excellence called for a *doubling* of Purdue's sales budget. Under McKinsey's prior tutelage, Purdue's promotional spending had already skyrocketed. McKinsey's ongoing influence on Purdue's operations after the 2007 guilty plea is stark:

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¹⁵⁸ Regarding the name change, CEO John Stewart wrote to McKinsey partners Rob Rosiello and Arnab Ghatak on August 15, 2013: "Paolo Costa was especially engaged in the discussion and he (among others) will be a champion for our moving forward with a comprehensive 'turbocharge' process – *though we do need to find a better and more permanently appropriate name*." PPLPC012000436626 (emphasis added).

\$50,000,000

\$45,000,000



All Promotional Spending on Non-Injectable Opioids by Defendant Purdue (Quarterly)

293. At the time of McKinsey's first known work for Purdue, Purdue spent approximately \$5 million per quarter on sales and marketing. By the time McKinsey's Project Turbocharge was implemented, total quarterly sales and marketing spending at Purdue exceeded \$45 million, an increase of 800%.

294. Project Turbocharge continued despite the arrival of a new CEO at Purdue. On January 17, 2014, new CEO Mark Timney received reports from McKinsey emphasizing that, in order to increase profits, Purdue must again increase the number of sales visits to "high-value" prescribers, i.e., those that prescribe the most OxyContin.¹⁵⁹

¹⁵⁹ In fact, recent deposition testimony suggests McKinsey may have been responsible for the fact that Timney was given the CEO job at Purdue in the first place. On October 30, 2020, Timney provided the following testimony (emphasis added):

Q: Are you familiar with McKinsey & Company?

A: I decline to answer on the ground that I may not be compelled to be a witness against myself in any proceeding.

Q: Did individuals at McKinsey assist you in getting hired as the CEO of Purdue?

A: I decline to answer on the ground that I may not be compelled to be a witness against myself in any proceeding.

295. Purdue and McKinsey worked together to implement "Turbocharging the Sales Engine."

296. McKinsey and Purdue also worked together on an "implementation plan" for E2E, with McKinsey taking on the role of "executive oversight" of projects including the creation of target lists, internal dashboards to track progress, and changes to Purdue's incentive compensation plan consistent with E2E.¹⁶¹

1. Targeting High Subscribers

- 297. Project Turbocharge called for revising the existing process for targeting high-prescribing physicians, with a shift from targeting solely on the basis of prescription deciles to considering additional factors. Based on its analysis, McKinsey told Purdue that "[t]here is significant opportunity to slow the decline of OxyContin by calling on more high-value physicians" and that "[t]he revenue upside from sales re-targeting and adherence could be up to \$250 million."
- 298. The core objective of McKinsey's initiative was to ensure that Purdue was "making calls on the highest potential customers with the right frequency to maximize prescribing potential."
- 299. McKinsey determined and advised Purdue that the top half of prescribing physicians "write on average 25 times more scripts per prescriber" than the lower half. McKinsey advised that Purdue would see a greater return on its sales investment by focusing on these targets, including on

In fact, McKinsey appears to have played a substantial role in the succession of several Purdue CEO's. Martin Elling, in his 2018 annual self-assessment, provided the following example of "how I deliver impact:" "Actively managing CEO/CXO transitions: ... from Michael Friedman to John Stewart (2007) to Mark Timney (2014) to Craig Landau (2017) at Purdue." MCK-MDL2996-0357931. "I drove our introduction of Purdue in 2004 and then, with Rob Rosiello, built it into a substantial and sustaining client... We have served across four CEOs and are now helping the new leadership team adapt to a world of headwinds for their core product OxyContin," he added. MCK-MDL2996-0357931.

¹⁶¹ MCK-MDL2996-0180338, at 0180340

prescribers with alarming prescribing patterns that raised red flags they were writing "prescriptions" for non-medical use. McKinsey's plan aimed at boosting sales of OxyContin by targeting the highest volume opioid prescribers, without addressing whether the expanded sales would be for an illicit market.

300. McKinsey found that Purdue did not "focus on the highest potential docs," measured both by the number of prescriptions and reimbursement considerations. ¹⁶² One McKinsey analyst urged McKinsey to recommend Purdue target "[1]iterally, at least all" prescribers in the top 20% of prescribers, "minus another few percent who are no sees[.]" McKinsey team lead Arnab Ghatak replied that "they probably have 20% no see[], but i'd also assume there are not many high writers that are no see." ¹⁶⁴ ("No see" prescribers are prescribers who do not accept visits from pharmaceutical sales representatives. Thus, McKinsey recognized that most of the highest volume prescribers, or "high writers" of prescriptions, were willing to entertain sales visits from sales representatives.)

301. "To put this in perspective," McKinsey stated,

the average prescriber in decile 5-10 [the top half of prescribers by volume] writes 25 times as many OxyContin scripts as a prescriber in decile 0-4. In Q1 2013 the majority (52%) of OxyContin primary calls were made to decile 0-4 prescribers. Including the secondary calls, 57% of the primary detail equivalents (PDEs) were made to decile 0-4 prescribers. Best practice in the industry is over 80% of effort on higher value prescribers."

- 302. McKinsey concluded: "Given that there are 14,000 uncalled physicians in deciles 5-10, there is significant opportunity to shift calls to higher potential prescribers." ¹⁶⁵
- 303. McKinsey pointed to a "true physician example" in Wareham, Massachusetts, who wrote 167 more OxyContin prescriptions after Purdue sales reps visited him. 166

¹⁶⁵ MCK-MDL2996-0187168 / PPLP004409892

¹⁶² MCK-MDL2996-0364024

¹⁶³ MCK-MDL2996-0364267

¹⁶⁴ Id

¹⁶⁶ PPLPC012000437356

True physician example Specialty Anesthesiology Location : Wareham, Massachusetts Market Decile: 8 12 months ending 12 months ending March 2012 March 2013 0 P1 18 P1 Calls made on physician 1 P2 1 P2 OxyContin scripts 177 344 written during 2nd half of year

Graphic from McKinsey presentation recommending targeting high prescribers

304. To slow or reverse the decline in OxyContin sales, McKinsey recommended a shift to "value deciles," which purported to weigh prescribers according to factors including overall opioid prescriptions, including the number of branded versus generic prescriptions; prescriber rules in place limiting sales calls; managed care access; and the number of the prescribers new to brand prescriptions, including new opioid patients and switches from other opioid products. The cumulative effect of the value rankings was to shift detailer emphasis onto the highest-volume prescribers. Further, McKinsey's analysis found that the highest-volume prescribers were themselves most influenced by detail visits.

305. Purdue moved quickly to do as McKinsey advised. All sales representatives received a memo on December 23, 2013, identifying how to select "SuperCore" prescribers, or the top ten targets, ¹⁶⁸ in their territory according to the E2E high prescribing principles and required that each SuperCore prescriber be visited at least twice a month. ¹⁶⁹

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¹⁶⁷ PPLPC022000646874

168 MCK-MDL2996-0316833

¹⁶⁹ PURCHI-000005915

1 170 2 As part of these changes, McKinsey's plan involved *more* minimum sales calls overall. ¹⁷¹ 3 306. 4 5 who later plead guilty to criminal charges related to an opioid drug ring. The 6 prescriber also surrendered his license to practice after an Ohio Medical Board investigation 7 revealed that he prescribed excessive and dangerous combinations of opioids and muscle relaxers 8 9 and that he prescribed opioids to a patient who complained of headaches and others who showed 10 signs of addiction. ¹⁷² The same prescriber received at least sixty visits from Purdue from mid-2013 11 through 2016.¹⁷³ 12 307. 13 14 The doctor, who worked at a family practice, was 15 charged with involuntary manslaughter, Medicaid fraud, drug trafficking, grand theft, and other 16 17 offenses. 18 308. 19 20 In 2018, the Drug Enforcement Administration issued an Order to Show Cause and 21 Immediate Suspension Order to Dr. Khan-Jaffrey over concerns that her DEA registration 22 "constituted an imminent danger to the public health and safety," finding she prescribed opioids 23 24 without a legitimate medical purpose and disregarded urine screens indicating abuse and 25 26 170 PPLPC022000686986 171 MCK-MDL2996-0187168 27 172 https://www.cnhinews.com/article 91ffac58-1b32-11e8-b264-6b34793bf5c3.html ¹⁷³ Public information about visits at which a payment was made is available for this time period through "Open 28 Payments."

¹⁷⁴ PPLPC014000257127

1	diversion. ¹⁷⁵ Dr. Khan-Jaffrey's DEA registration was fully revoked on July 28, 2020. ¹⁷⁶ Dr. Louis
2	Spagnoletti, of Marlton, New Jersey, a
3	lost his state license to prescribe controlled substances in 2018. Similarly, Dr. Vivienne
4	Matalon, a Decile 10 prescriber from Cherry Hill, New Jersey,
5	went on to lose her license in 2018 as well, for allegedly receiving kickbacks to
6	prescribe the fentanyl drug Subsys to three patients, including one that died. 180
7	309. Another prescriber, Dr. Damon Cary of Wilmington, Delaware,
8 9	
9	¹⁸¹ received an emergency
10	suspension order in 2019 after prescribing controlled substances, including opioids, to undercover
11	officers without performing any medical examinations. ¹⁸² Dr. Eva Dickinsson, of Harrington,
12 13	Delaware,
13 14	was arrested on marijuana charges in 2016 and had her license suspended in 2017 for
15	sharing drugs, including opioids, with her patients. 184
16	310. Dr. Michael Cozzi of Fort Wayne, Indiana
17	had his medical license suspended
18	in 2016, where he had prescribed more controlled substances than any other Indiana prescriber,
19	
20	with over two million doses of oxycodone, seeing ninety to 100 patients a day. 185 Dr. Jamie Gurrero,
21	
22	https://www.federalregister.gov/documents/2020/07/29/2020-16387/kaniz-f-khan-jaffery-md-decision-and-order
23	¹⁷⁶ Id. ¹⁷⁷ PPLPC014000257127
24	https://patch.com/new-jersey/moorestown/state-suspends-doctor-accused-illegally-prescribing-opioids PPLPC014000257127
25	¹⁸⁰ https://nj.gov/oag/newsreleases18/pr20180504d.html ¹⁸¹ PPLPC014000257130
26	 https://www.delawareonline.com/story/news/health/2019/08/05/doctor-prescribed-opioids-undercover-cops-failed-follow-protocol/1920386001/ PPLPC014000257130
27	¹⁸⁴ https://www.delawareonline.com/story/news/health/2017/01/19/doctors-license-suspended-delaware-
28	maryland/96779080/ 185 https://www.wane.com/news/fort-wayne-pain-doctors-medical-license-suspended/ (He later died in a tractor accident, https://www.journalgazette.net/news/local/police-fire/20180816/tractor-accident-kills-pain-doctor)

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was sentenced to 100 months in prison in 2016 after pleading guilty to unlawful distribution or dispensing of controlled substances, health care fraud, conspiracy, and money laundering.¹⁸⁷

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Misrepresentations to these prescribers were especially insidious because they were aimed at general practitioners who lack the time and expertise to closely manage higher-risk patients on opioids.

312. McKinsey also urged, consistent with continually refining its granular approach, that sales representatives devote two-thirds of their time to selling OxyContin and one-third of their time selling Butrans, another Purdue opioid product. Previously, the split had been fifty-fifty.

2. Circumventing Safeguards Against Abuse and Diversion

- 313. Project Turbocharge also involved a granular analysis of Purdue's individual sales channels. In its August 8, 2013, report to the Purdue board, McKinsey also attributed the decline in OxyContin sales to safeguards to limit suspicious opioid sales. McKinsey informed Purdue that "[t]he retail channel, both pharmacies and distributors, is under intense scrutiny and direct risk." "There are reports of wholesalers stopping shipments entirely to an increasing number of pharmacies," "[m]any wholesalers are also imposing hard quantity limits on orders based on prior purchase levels," and "[p]harmacy chains are implementing guidelines for which patients can fill opioid prescriptions[.]" 188
- 314. For instance, McKinsey recommended that Purdue circumvent pharmacies entirely with a mail order program because enforcement by federal regulators was decreasing OxyContin

¹⁸⁶ PPLPC014000257130

 $^{^{187}\} https://www.justice.gov/usao-wdky/pr/kentuckiana-anesthesiologist-sentenced-100-months-unlawful-distribution-controlled$

¹⁸⁸ MCK-MAAG-0024297

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¹⁸⁹ *Id*.

¹⁹⁰ MCK-MDL2996-0290827. 27

¹⁹¹ MCK-MDL2996-0041646.

¹⁹² MCK-MDL2996-0104431; MCK-MDL2996-0041646.

¹⁹³ PPLPC012000437346.

¹⁹⁴ *Id*.

dispensing through Walgreens. McKinsey informed the Sacklers that "[d]eep examination of Purdue's available pharmacy purchasing data shows that Walgreens has reduced its units by 18%." Further, "the Walgreens data also shows significant impact on higher OxyContin dosages." ¹⁸⁹

315. In order to counter these perceived problems, McKinsey suggested that Purdue's owners lobby Walgreens specifically to increase sales and circumvent the safeguarding sales limits. It also suggested the establishment of a direct-mail specialty pharmacy so that Purdue could circumvent Walgreens and sell directly to Walgreens' customers. Finally, McKinsey suggested the use of opioid savings cards distributed in neighborhoods with Walgreens locations to encourage the use of Purdue's opioids despite Walgreens actions.

316. McKinsey's initiative also included ways to circumvent these safeguards. McKinsey recommended that the sales force distribute vouchers and "starter kits" for patients who faced copays for OxyContin prescriptions. 190 In particular, McKinsey recommended dispensing vouchers to outlets of a specific large national pharmacy chain where prescriptions and OxyContin inventories were down. 191 This chain, as part of its own settlement with the Drug Enforcement Administration, had removed pharmacist bonuses for dispensing opioids. 192

3. Incentivizing Opioid Sales

317. McKinsey's "turbocharging" plan also had other elements.

ix. McKinsey's Efforts Triple OxyContin Sales

- 318. In 2013, despite significant headwinds, with marketing activities turbocharged, OxyContin sales peaked. The restrictions on Purdue's sales and marketing methods contained in the Corporate Integrity Agreement should have resulted in fewer overall OxyContin sales; the guilty plea identified a specific segment of existing OxyContin sales that were illegitimate and should thus cease. All else being equal, OxyContin sales should have decreased to account for the successful elimination of improper sales. In fact, OxyContin sales did decrease in the immediate aftermath of the 2007 guilty plea.
- 319. And within five years, however, OxyContin sales would triple. McKinsey is responsible for the strategy that accomplished this. It presented specific plans to Purdue, which Purdue adopted and spent hundreds of millions of dollars implementing. The result: a final spasm of OxyContin sales before the inevitable decline of the drug.¹⁹⁵
- 320. The Purdue McKinsey collaboration was a spectacular success. Between the 2008 and 2016, Purdue distributed in excess of \$4 billion to the Sackler family, with \$877 million distributed in 2010 alone.
- 321. These distributions would not have been possible without the McKinsey's work dramatically increasing OxyContin sales.
- 322. The Sacklers were aware of the value McKinsey provided: on December 2, 2013, CEO John Stewart informed Kathe Sackler and Vice President of Sales and Marketing Russell Gasdia Project Turbocharge "was already increasing prescriptions and revenue." Crucially, these results were already being realized *before* the strategy was fully deployed as the theme of the 2014 National Sales Meeting. Stewart elaborated to Sackler that "trends are more positive than was the

¹⁹⁵ On February 10, 2018, Purdue announced that it is no longer marketing opioids, and disbanded its OxyContin sales force.

case a few months back, and when the E2E Project (the changes arising out of the McKinsey analysis) is fully implemented there will certainly be additional increases." ¹⁹⁶

323. Later that month,

- 324. McKinsey's contributions to Purdue's growth after 2007 are remarkable. OxyContin sales should have naturally declined; the Department of Justice identified OxyContin sales that were illegitimate because of Purdue's conduct, and the Inspector General of the Department of Health and Human Services entered into a Corporate Integrity Agreement whereby Purdue was monitored to assure that those sales did not continue.
- 325. In 2007, the year of Purdue's guilty plea, net sales of OxyContin totaled approximately \$1 billion. 198
- 326. The guilty plea "did little to stem Purdue's blistering growth rate." In fact, by 2010, after McKinsey was advising Purdue on how to maximize sales, OxyContin sales exceeded \$3 billion: a *tripling* of revenue from OxyContin sales.¹⁹⁹
- 327. Under McKinsey's guidance, OxyContin sales would reach their all-time peak in 2013, the year McKinsey proposed, and Purdue adopted, Project Turbocharge.²⁰⁰ That OxyContin sales peaked in 2013 is especially notable, given that *overall* opioid prescriptions had *already peaked* three years earlier, in 2010.²⁰¹ McKinsey's efforts added a final boost to OxyContin sales before the eventual unraveling, and Purdue's decision, in the end, to cease marketing the drug.

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¹⁹⁶ PPLPC012000454422

¹⁹⁷ PPLPC012000457292

¹⁹⁸ See David Crow, How Purdue's 'one-two' punch fueled the market for opioids, Financial Times, September 9, 2018, available at: https://www.ft.com/content/8e64ec9c-b133-11e8-8d14-6f049d06439c

²⁰⁰ Phil McCausland and Tracy Connor, *OxyContin maker Purdue to stop promoting opioids in light of epidemic*, NBC News, February 10, 2018, *available at:* https://www.nbcnews.com/storyline/americas-heroin-epidemic/oxycontin-maker-purdue-stop-promoting-opioids-light-epidemic-n846726

²⁰¹ Gery P. Guy Jr, *at al.*, *Vital Signs: Changes in Opioid Prescribing Patterns in the United States*, 2006-2015, Centers for Disease Control and Prevention, July 7, 29017, *available at:* https://www.cdc.gov/mmwr/volumes/66/wr/mm6626a4.htm

²⁰⁴ PPLPC029000547371

332. Part of the unique value McKinsey provides is its deep knowledge of its clients' competitors, often because it counts those same competitors as its clients. McKinsey generally does not disclose to its clients its work for their competitors.

333. The opioid industry was no different. Indeed, McKinsey specifically worked to

i. Endo

334. While McKinsey was working for Purdue, McKinsey was also working for Endo Pharmaceuticals. Arnab Ghatak was a principal McKinsey partner on both accounts at the same time. There was additional overlap between the McKinsey teams staffed to Purdue and Endo, including McKinsey partners Nicholas Mills and Laura Moran. After all, these particular consultants had granular expertise in the specific subject-matter relevant to these opioid manufacturers. That subject-matter expertise is a compelling reason why McKinsey is hired in the first place. McKinsey advised both Endo and Purdue how to maximize the sales or their branded opioid products—Belbuca (Buprenorphine), Butrans (Buprenorphine), Opana (Oxymorphone), and OxyContin (Oxycodone) —all at once.

1. New Blues

335. Like Purdue, Endo was historically a pharmaceutical manufacturer focused on the pain market. Like Purdue, Endo relied on opioid sales for a significant portion of its business. As a

²⁰⁵ MCK-MDL2996-0041741.

Ghatak's familiarity with both Endo and Purdue is perhaps one reason why, on April 3, 2014, Ghatak was placed in charge of analyzing a proposed *partnership* between Purdue and Endo to sell opioids. Lauran Moran described the "partnership workstream" that McKinsey was then performing for Purdue to identify ways for Purdue to obtain near-term growth. She stated that Purdue and McKinsey "agreed the partnerships workstream should include the top 3 potential partners (Valeant, Endo and Pfizer for now). And for each what assets each partner would bring and what growth (most importantly) would the deal bring. Arnie [Ghatak] and Phil to work out Endo and Valeant, John G and Raul to do Pfizer tomo." MCK-MDL2996-0421790.

matter of fact, Endo's history with opioids predates the Sacklers' ownership of Purdue. In 1950, Endo's predecessor, Intravenous Products of America, Inc., launched Percodan, an Oxycodone/Aspirin tablet. In 1971, Endo, then owned by E.I. du Pont de Nemours and Company ("DuPont"), launched Percocet, another oxycodone-based tablet.²⁰⁷

- 336. In 1997, Endo separated from DuPont to become a standalone private company retaining Percodan and Percocet.²⁰⁸ In 2000, as the result of an acquisition, the company became public.²⁰⁹
- In 2006, Endo launched its own branded oxymorphone products, Opana and Opana ER. 210 With the legacy assets of Percodan and Percocet, Endo's business had always been focused on opioid sales. Oxymorphone is not a new opioid, and Opana was not Endo's first oxymorphone product. It was first synthesized more than a century ago in Germany. Endo began selling it in the United States in 1959 under the name Numorphan.
- 338. Numorphan was referred to as "blues," after the color of the 10mg pills. It delivered a more euphoric high than heroin, according to some. In 1974, the National Institute on Drug Abuse noted in its "Drugs and Addict Lifestyle" report that Numorphan was popular as an abused drug for its quick and sustained effect.²¹¹ By 1979, Endo withdrew Numorphan from the market. Upon information and belief,

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²⁰⁷ https://www.endo.com/about-us/history#fragment-25 ²⁰⁸ https://www.endo.com/about-us/history#fragment-24

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²⁰⁹ https://www.endo.com/about-us/history#fragment-21 ²¹⁰ https://www.endo.com/about-us/history#fragment-15

²¹¹ John Fauber & Kristina Fiore, Abandoned Painkiller Makes a Comeback, MedPage Today (May 10, 2015), available at: https://www.medpagetoday.com/psychiatry/addictions/51448

²¹² EPI000443330 ("

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ENDO-OPIOID-MDL-06246554

339. The memory of the 1970s Numorphan addiction crises did not fade quickly. In 1989, the film *Drugstore Cowboy* featured Matt Dillon as an addict in the 1970s who robs drug stores to obtain drugs to sell in order to finance his opioid dependency.²¹³ In one scene, an addict asks Dillon's character if he has any "blues." Dillon's character explains that "blues" are increasingly hard to find, and offers to sell morphine sulfate to an addict instead. The addict explained that he much preferred the Numorphan, but settled for the morphine.²¹⁴

- 340. With the launch of Opana, Endo decided it was time for history to repeat itself. After Opana's approval in 2006, Endo solidified its position as a pain specialist among manufacturers. By 2012, opioid sales accounted for approximately \$403 million of Endo's \$3 billion in revenue, more than 10%. From 2010 to 2013, total Opana ER revenue alone exceeded \$1.1 billion.
- 341. Opana and Numorphan were both oxymorphone. The brand name was the only thing that changed. What Endo removed from the market in 1979 due to abuse concerns, it re-introduced 27 years later. After 2006, Opana was on occasion referred to as "blue heaven," or, more to the point, "new blues."
- 342. In 2017, Endo would once again remove its branded oxymorphone product from the market, and for the same reason. Endo's abuse-deterrent formulation of Opana was removed at the request of the FDA due to acute concerns about its abuse potential.
- 343. In addition to its branded products, Endo, through subsidiaries Qualitest Pharmaceuticals, Inc. and, after its acquisition in 2015, Par Pharmaceuticals, also manufactured generic versions of oxycodone, oxymorphone, hydromorphone, and hydrocodone. Over the course of McKinsey's relationship with Endo, McKinsey would repeatedly advise Endo how to maximize its generics business in addition to sales of Endo's branded opioids.

²¹³ See https://www.imdb.com/title/tt0097240/; https://www.bionity.com/en/encyclopedia/Oxymorphone.html

²¹⁴ Scene from Drugstore Cowboy, available at: https://www.youtube.com/watch?v=TksvZdrx9_A

²¹⁵ https://www.deadiversion.usdoj.gov/drug_chem_info/oxymorphone.pdf

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2. **Old Friends**

- 344. McKinsey's relationship with Endo began as early as in 2006, the same year as the Opana launch.
- 345. McKinsey's earliest known work with Endo concerned the launch of Opana in Europe, but its relationship with Endo would expand to encompass all aspects of Endo's business, including corporate organization and resource allocation, the launch of a new branded Buprenorphine product, and sales force optimization efforts for Endo's branded and generic opioid products.
- In 2007, McKinsey was shaping overall corporate strategy at Endo. In a presentation 346. to Endo's board of directors in May of that year

- 347. McKinsey's partnership with Endo would last more than a decade, and, like its relationship with Purdue, is an exemplary example of the transformational relationship in action.
- 348. In some ways, the McKinsey's relationship with Endo was even more tightknit and companionable than with Purdue. For instance, no one at Purdue previously worked for McKinsey. In early 2013, Rajiv de Silva, previously a leader of McKinsey's PMP group, was appointed CEO of Endo. At Endo, McKinsey was now advising an old friend, one of its previous senior partners.²¹⁷

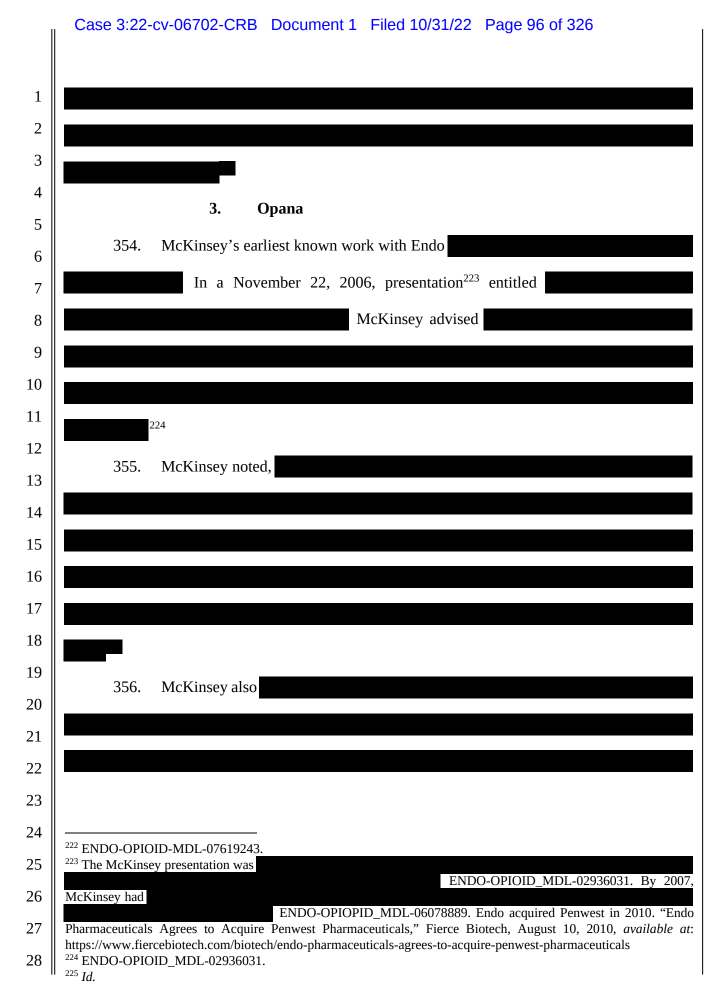
²¹⁶ ENDO-OPIOID_MDL-02899510.

²¹⁷ See "Rajiv De Silva Named President and CEO of Endo Health Solutions," Press Release dated February 25, 2013, available at: https://investor.endo.com/news-releases/news-release-details/rajiv-de-silva-named-president-and-ceoendo-health-solutions ("Earlier in his career, he was a Principal at McKinsey & Company, where he served as a member of the partnership group that led the global Pharmaceuticals and Medical Products practice.")

1 As de Silva himself explained, 349. 2 3 4 Under de Silva, Endo relied more heavily on McKinsey than ever before. McKinsey 350. 5 consultants interacted directly and often exclusively with de Silva. McKinsey was so close to the 6 Endo CEO that it could intervene in direct reporting from one of de Silva's deputies.²¹⁹ It is as if 7 8 McKinsey had insinuated itself as a shadow layer of bureaucracy within Endo. 9 351. McKinsey maintained weekly performance review meetings with de Silva and 10 senior Endo management. In these meetings, granular weekly sales data was reviewed for each of 11 Endo's branded products, including Opana.²²⁰ 12 352. McKinsey advised both Purdue and Endo contemporaneously for more than a 13 decade. With each client, the goal was the same: to maximize opioid sales. The work McKinsey 14 performed for each client was so similar that there was routinely confusion internally about whether 15 a specific project or task to perform was for Endo or Purdue.²²¹ 16 17 353. Despite McKinsey's emphasis on confidentiality, the fact that McKinsey repeats its 18 work from one client to the next is well-known to the client. Indeed, it is part of the justification in 19 hiring McKinsey in the first place. McKinsey can tell you what everyone else is doing. 20 21 22 23 24 ²¹⁸ ENDO-OPIOID-DEPMAT-000047877 at pg. 320:22 – 321:3. 25 ²¹⁹ See MCK-MDL2996-0405502 (Email from Ghatak to de Silva, stating that it "would be great for you to push Blaine and Bob [both Endo employees] on why there are no slides showing the metrics on field call attainment . . . there was 26 an explicit agreement to track them. Setting the expectation that you want them included would really help"). ²²⁰ MCK-MDL2996-0062712. 27 ²²¹ In response to an internal email from Craig MacKenzie to other McKinsey consultants seeking "expert input on labels for abuse deterrent formulations" in conjunction with McKinsey's work on the Belbuca launch (discussed infra.), 28 McKinsey consultant Jeff Smith replied, "Craig - is this for Purdue or Endo? If for Endo, I am conflicted." MCK-

MDL2996-0383805.

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15	359. Within a few years of its introduction in the United States, abuse of the drug became
16	widespread. Endo then sought to introduce a reformulated version of Opana that it could market as
17	abuse-deterrent by introducing a tamper-resistant coating to the pill.
18	360. In December 2011, Endo obtained FDA approval for a new formulation of Opana
19	ER with the coating that Endo claimed was crush-resistant. The following month, however, the
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21	FDA told Endo that it could not market Opana ER, even after the reformulation, as abuse-deterrent.
22	361. Endo "did not submit any new clinical safety or efficacy data" as part of its
23	application, but rather relied entirely on the "bioequivalence" of the new and old formulations of
24	Opana. Obtaining approval of reformulated Opana ER on this basis allowed Endo to rely on the
25	safety and efficacy of the original version of the drug as the basis for approval of the reformulated
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28	²²⁶ Id. ²²⁷ Id.
I	²²⁸ Id.

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version.²²⁹ The FDA found that such promotional claims "may provide a false sense of security since the product may be chewed and ground for subsequent abuse." In other words, Opana ER was still crushable. In December 2011, Endo admitted that "[i]t has not been established that this new formulation of Opana ER is less subject to misuse, abuse, diversion, overdose, or addiction."²³⁰

362. In 2013, an Endo training module directed key opinion leaders to instruct prescribers that OPANA ER with INTAC is the only oxymorphone designed to be "crush-resistant," and advised the key opinion leaders to state during their speeches that "[t]he only way for your patients to receive oxymorphone ER in a formulation designed to be crush-resistant is to prescribe OPANA ER with INTAC."231 The speakers were advised to stress that generic versions of Oxymorphone "are not designed to be crush-resistant."

363. These abuse-deterrent attributes of the reformulation—the very characteristics McKinsey and Endo touted as a reason to prescribe Opana—were a sham. The reformulation was designed to prevent the pill from being crushed and snorted through the nose. It did not prevent intravenous use, however. The result was that many users already dependent of Opana began using needles to inject the drug for the first time. As an internal Endo email put it,

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Jeff, a veteran of the war in Iraq, explained the process. Jeff first became dependent 364. on Percocet and Opana after returning from Iraq, where his back was injured when his Humvee rolled over in 2008. After being prescribed opioids for his back pain, Jeff became dependent, and

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²²⁹ Intervenor Impax Laboratories, Inc.'s (1) Cross-Motion to Dismiss; or, in the Alternative, (2) Opposition to Plaintiff's Motion for a Preliminary Injunction, Endo Pharmaceuticals Inc. v. U.S. Food and Drug Administration, et al. ("Impax Br."), No. 1:12-cv-01936 Doc. 18 at 7 (D.D.C. Dec.9, 2012); see also FDA Summary Review for Regulatory Action, NDA 201655 (Dec. 9, 2011) (stating that "[n]o new safety data were included in this submission" and "[n]o efficacy studies were submitted in this application.").

²⁷ ²³⁰ Endo Dec. 12, 2011 News Release; Ex. A to Rurka Decl., Endo Pharmaceuticals Inc. v. U.S. Food and Drug Administration, et al., No. 1:12-cv-01936 Doc. 18-2 (D.D.C. Dec. 9, 2012). 28

²³¹ EPI000421543.

²³² END00010732.

began using Opana by snorting it. Endo then introduced the reformulated abuse-deterrent version of Opana in 2012. "[A]nd then they reformulated them," he said, referring to the Opana pills. "And the only way you could really do them is inject them because if you actually swallow them, it – you – they really don't do nothing."²³³

365. Jeff and his companion Joy showed the journalist how the drug was used. "You want to see how to cook it?" Jeff asked. He and Joy then proceeded to place a portion of an Opana pill on piece of aluminum and heat it with a lighter. "Right away, I can start to see this hard, white coating just kind of floating off the piece of the pill. It looks like plastic," described the journalist witnessing the process.²³⁴

366. Joy explained how the abuse-deterrent coating, once melted, was discarded by using the filter of a cigarette: "Now you see the coating of – all that mess laying there still? . . . That's what the filter's for." The journalist described what then took place:

And Joy puts that cigarette filter into the liquid, and they Joy, Jeff and another guy each take turns with their needles, sticking it into the filter and pulling the liquid through. Joy and Jeff turn their back to me while they inject. And then it just gets really, really quiet.

367. Joy couldn't conceive of the position she was in. A nurse, she hurt her back at work and began taking prescription pain medication. She began taking the pills by mouth, and later began to snort them. Dependency began. But she told herself, "I'd never ever would use a needle, never. I'm never going to do that."²³⁶

²³³ Kelly McEvers, "Opioid Epidemic Sparks HIV Outbreak in Tiny Indiana Town," NPR, March 31, 2016, *available at*: https://www.npr.org/2016/03/31/472577254/opioid-epidemic-sparks-hiv-outbreak-in-tiny-indiana-town

 $^{^{234}}$ *Id*.

²³⁵ *Id*.

²³⁶ *Id*.

1 2 3 4 5 373. McKinsey indicated 6 7 8 9 10 11 12 In addition to being neither feasible nor safe/ethical, the study was beside the point. 374. 13 An insufflation study is meant to determine the abuse characteristics of a drug when used nasally— 14 i.e., by snorting the drug.²⁴⁴ The relevant concern for Opana's reformulated version was *injection*, 15 not insufflation. 16 17 But the insufflation study worked for Purdue. Going forward, McKinsey suggested 375. 18 19 As the preceding paragraphs make clear, Endo and McKinsey were laser-focused 376. 20 on maximizing overall sales of Endo products, and decidedly not on concerns over their actual 21 abuse potential or the appropriate size of the market for these products, given evident, longstanding, 22 and ever-present concerns about their abuse. To the point, McKinsey regarded concerns about 23 24 25 ²⁴¹ EPI002107711. ²⁴² *Id*. 26 ²⁴³ *Id.* (emphasis added). ²⁴⁴ See General Principles for Evaluating the Abuse Deterrence of Generic Solid Oral Opioid Drug Products, FDA 27 Drug Evaluation and Research, November 2017, available at: https://www.fda.gov/files/drugs/published/General-Principles-for-Evaluating-the-Abuse-Deterrence-of-Generic-28 Solid-Oral-Opioid-Drug-Products-Guidance-for-Industry.pdf ²⁴⁵ EPI002107711.

opioid abuse only as a means by which its clients could introduce *differentiated* products (i.e., those with purported abuse-deterrent or tamper-resistant features) to continually perpetuate overall opioids sales for their clients. In all instances, the parties desired for the size of that overall opioids market to grow in line with the introduction of "differentiated" products like a reformulated Opana.

377. Endo's purported concern about deterring abuse of its drugs was laid bare as farce by a particularly striking decision: to continue to sell the old formulation of Opana despite touting the notion that the old formulation was purportedly dangerous in ways that the new formulation was not. Endo

378. Endo not only continued to distribute original Opana for nine months after the reformulated version became available, it declined to recall original Opana ER despite its dangers.²⁴⁷ In fact, Endo also claimed in September 2012 to be "proud" that "almost all remaining inventory" of the original Opana ER had "been utilized."²⁴⁸

379. In June 2013, an Endo employee informed the McKinsey consultants of a

4. Belbuca: Endo's Answer to Butrans

380. Buprenorphine is another differentiated product. Opioid manufacturers began to introduce Buprenorphine products to the market after the introduction of OxyContin, Opana, and

²⁴⁶ ENDO-OPIOID-MDL-02324795.

²⁴⁷ Impax Br. at 1.

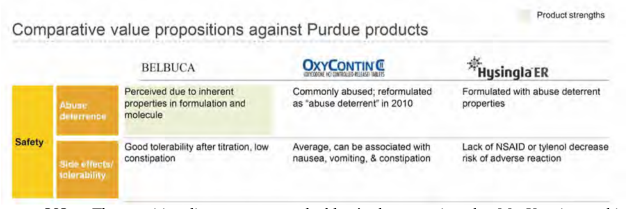
²⁴⁸ *Id.*; Endo News Release, Sept. 6, 2012 (Ex. L to Rurka Decl) *Endo Pharmaceuticals Inc. v. U.S. Food and Drug Administration, et al..*, No. 1:12-cv-01936 (Doc. 18-4) (D.D.C. Dec. 9, 2012).

²⁴⁹ ENDO-OR-CID-00400235 (emphasis added).

other branded opioids long-known to have abuse and dependency problems. Buprenorphine products were marketed as purportedly less dangerous than products such as OxyContin or Opana.

381. Of course, Endo and Purdue continued to assiduously market and sell OxyContin and Opana alongside their Buprenorphine products, and McKinsey worked with each at every step of the way, despite the implicit contradiction in marketing two products at the same time whose point of differentiation is one being *less addictive and dangerous* than the other.

382. For example, on August 13, 2015, McKinsey's Craig MacKenzie circulated a discussion document to Endo and McKinsey staff entitled "Belbuca value proposition," which laid out McKinsey's thoughts on how to differentiate Endo's buprenorphine product from other opioids in the marketplace. One point of differentiation McKinsey noted was that OxyContin was commonly abused, while Endo's Belbuca hopefully would not be: 251



383. The cognitive dissonance was palpable. At the same time that MacKenzie sent his email differentiating Belbuca, and as described *supra*, McKinsey was *also* maximizing OxyContin sales for Purdue—the opioid it was describing to Endo as commonly abused.

²⁵⁰ MCK-MDL2996-0410742.

²⁵¹ MCK-MDL2996-0006669, at 0006675.

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Butrans was Purdue's buprenorphine product.

385. Once Butrans was launched at Purdue, McKinsey worked with Endo to create another branded Buprenorphine product to compete with Butrans. These product planning and launch processes are long-term affairs. McKinsey worked with Endo on this project for *four years* before Endo's Belbuca obtained FDA approval.

386. McKinsey remained in place at Endo to implement the launch of Endo's Buprenorphine product. The strategic goal of Belbuca—the key to its commercial success—was to convert short acting opioid ("SAO") users to Belbuca. As McKinsey explained to CEO Rajiv de Silva, "The fundamental question is whether Belbuca will take share from the short-acting opioids."²⁵²

387. Ultimately, Belbuca was not a large commercial success for Endo because it failed to transition a sufficient number of short acting opioid users to the long-acting Belbuca. As the drug underperformed, Endo felt ever more pressure to stimulate sales. John Harlow described one meeting with de Silva on April 8, 2016: "We just got out of the review with Rajiv and clearly our TRx trends are not good and are behind other recently launched pain products . . . the request from Rajiv was to do anything possible that could be implemented ASAP to stimulate RXs." ²⁵³

388. By July 6, 2016, McKinsey and Endo were increasingly focused on converting short-acting opioid users.

Notably, the goal was to increase *overall* buprenorphine prescriptions,

²⁵² MCK-MDL2996-0210158.

²⁵³ MCK-MDL2996-0358973, at 0358975.

²⁵⁴ ENDO-OPIOID_MDL-07264539

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not only those of Belbuca. The discussion document identified an objective to create "a new treatment paradigm for [Buprenorphine] and Belbuca at the transition between SAO and LAO."255 In order to do so, the discussion group needed to determine "what medical support we need to position Buprenorphine as the best transition from SAO to LAO."256

389. McKinsey provided a slide to Endo describing the way that narrative would first be created and then exploited for market positioning:²⁵⁷

²⁵⁵ *Id*.

²⁵⁷ MCK-MDL2996-0382731.

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Overall timeline for rolling out Belbuca/Bup 'transition' positioning Q3 - Get the facts in place to tell the story Q4 - Broadcast the story 1H 2017 - Create compelling data to accelerate the proof points Medical Literature Publish national article Make progress on RWE/RCT Review on Bup transition trials (e.g., H2H with Oxy, cancer pain) Crafting the overall Build out speakers on cross-stakeholder story Bup Finalize retrospective studies for Bup, as ideal on abuse Drive 'can't step through 'transition' CII' message to payers Health Economics analysis Prioritize stakeholders Create pilots with Exploration of 'step through (payers, advocacy payers/providers (e.g., Bup' deals with payers (prior groups) for VA, Geisinger) to prove to CII LAO) communication value This report is sicilely for the use of client personnel. No part of it may be endo , circuisled, guoled, or reproduced for distribution outside the client organization without prior written approval from Legal

390. McKinsey and Endo referred to this effort to revive Belbuca sales by promoting buprenorphine as a bridge to long-acting opioid use as a "moonshot." One aspect of this "moonshot" would be that Belbuca (and Buprenorphine, generally) would convert short-acting opioid users to long-acting opioids users of products *other than Buprenorphine*. McKinsey and Endo instead conceived of Belbuca as an "*Initial* ATC Opioid Therapy." It was to be positioned as "the *first* LAO for poorly controlled or dissatisfied chronic pain patients transitioning from short-

²⁵⁸ MCK-MDL2996-0382412.

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- 391. Thus, the overall marketing strategy McKinsey assisted Endo in designing and deploying for Belbuca was designed to transition ever more patients to long-acting opioids. Belbuca could find its market niche as a stepping stone as individuals proceed through the patient funnel from short acting opioid users to longer-term long-acting opioid users. The farther individual proceeds through this funnel, the more the individual is worth.
- 392. McKinsey also knew that this same pathway that begins with opioid therapy after a serious injury also leads to opioid dependency and addiction. In 2011, McKinsey was working on

²⁵⁹ MCK-MDL2996-0404374, at 0404375.

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"Project X." which was the project to develop a buprenorphine product to compete with Butrans. (Belbuca, in other words, was the result of Project X.) McKinsey described the "opioid dependence treatment pathway" as follows: 260

OPIOID DEPENDENCE TREATMENT PATHWAY

- Patient begins opioid therapy following serious injury/surgery
- Patient pain not controlled, dosages increase, patient seeks additional medication options
- Patient referred to pain specialist
- Patient shows up in ER, Clinic or seeks out addictionologist
- Addictionologist Pain Specialist or Psychiatrist with additional license to prescribe Methadone, Subutex/Suboxone for opioid dependence
- Options
 - Methdone
 - Subutex initiation, followed by Suboxone treatment

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393. In the same presentation, McKinsey identified the key to a successful launch of a branded Buprenorphine product: "The challenge faced by Endo will not be to gain formulary approval, it will be to gain tier 2 status and *minimize restrictions on prescribing*." ²⁶¹

5. Turbocharging the Sales Force with a Blitz

394. In 2015, a McKinsey team led by Arnab Ghatak proposed to Endo a sales transformation to invigorate Endo's product sales, including its opioids. At the suggestion of

²⁶⁰ MCK-MDL2996-0131500, at 0131512.

²⁶¹ MCK-MDL2996-0131500, 0131525 (emphasis added).

1 Ghatak, McKinsey used the Purdue Pharma Project Turbocharge model from the previous year as 2 a template for the Endo proposal.²⁶² Even the PowerPoint presentations used to create the proposal 3 to Endo were drafted off the Project Turbocharge slides. On June 28, 2015, Sherin Ijaz of McKinsey 4 emailed Ghatak, Nicholas Mills, and Laura Moran to circulate a draft proposal for an "Endo sales 5 force transformation" PowerPoint presentation. Ijaz explained, "Laura, I heavily leveraged what 6 you send (sic) from Purdue as it was all applicable." ²⁶³ All three of the recipients of Ijaz's email 7 regarding the Endo proposal had been working on the Purdue account for years. 8 9 395. Endo 10 11 12 13 396. Endo's Vice President & General Manager of its Pain Business Unit, John Harlow, 14 15 16 17 18 19 20 397. The Endo and Purdue proposals were essentially identical sales transformations. The 21 goals were the same: to maximize sales of opioids. Merely the names were changed. While 22 McKinsey offered to "turbocharge" Purdue's sales force, McKinsey proposed a "sales force blitz" 23 24 for Endo.²⁶⁷ 25 26 ²⁶² MCK-MDL2996-0075895. ²⁶³ MCK-MDL2996-0070237.

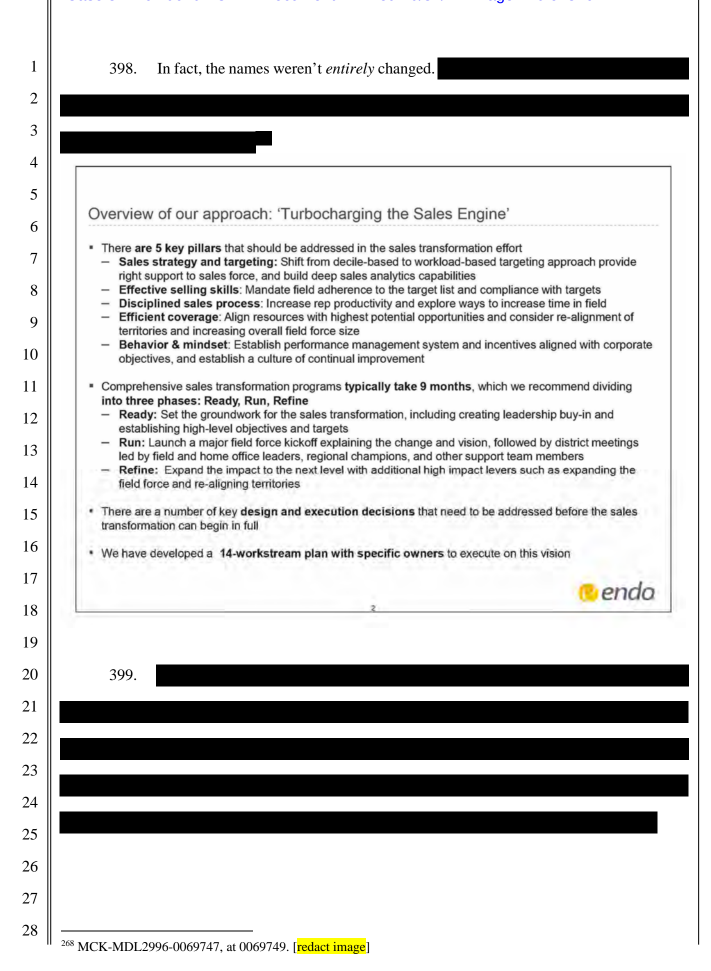
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²⁶⁵ *Id*.

²⁶⁴ ENDO AAC 00363406.

²⁶⁷ E.g., MCK-MDL2996-0130803; MCK-MDL2996-0132851.



1 400. 2 3 4 5 401. Upon McKinsey's suggestion, Endo began reallocating sales force resources to 6 Opana from other Endo products such as Sumavel, a migraine medication, and Voltaren, an anti-7 inflammatory.²⁶⁹ Writing to the McKinsey team, Endo's Alicia Logan stated the joint mission, "I 8 9 agree that our main goal is to maximize the increased promotional efforts for [Opana ER] without 10 disrupting/sacrificing [Sumavel] or [Voltaren] TRx volume and it appears that we [can] accomplish 11 this with your recommendation of addition another 500 targets."270 12 402. With the Sales Force Blitz underway, Endo received good news in New York. Years 13 prior, Endo had initiated patent litigation against generic manufacturers of Opana ER, arguing that 14 the generic versions of the drug infringed on Endo's patents. In part because of the perceived 15 16 impending loss of exclusivity, Endo had in recent years allocated its sales force capacity away from 17 Opana and to other Endo products. 18 403. On August 14, 2015, Endo received a favorable initial ruling declaring that the 19 generic versions of Opana violated Endo's patents, and enjoined their further sale. The ruling 20 provided additional patent exclusivity for Opana, and Endo was keen to exploit its advantage. 21 404. That afternoon, 22 23 24 25 26 27 ²⁶⁹ MCK-MDL2996-0409466. 28 ²⁷⁰ MCK-MDL2996-0409436, at 0409437 (sic).

²⁷¹ ENDO-OPIOID MDL-02279530.

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405. The following week, Harlow wrote to the McKinsey team working for Endo to focus their attention on Opana ER. "Now with our litigation victory from last week, plus our UHC opportunity, there is an increased need to increase FF support to drive Sep-Dec business. . . . With this win, I am now willing to go broader with OER targeting."²⁷²

406. McKinsey and Endo proceeded to design and implement retargeting strategies to boost Opana sales in late 2015.

ii. Johnson & Johnson/Janssen

407. McKinsey also working with Johnson & Johnson, whose role overseeing and contributing to the opioid crisis has been exhaustively detailed in other complaints. *See, e.g., City and County of San Francisco v. Purdue Pharma L.P.*, N.D. Cal. No. 18-2591, Doc. 128 (Mar. 13, 2020). Johnson & Johnson occupied multiple roles within the opioids industry. Through its subsidiary, Janssen Pharmaceuticals ("Janssen"), it marketed and sold branded opioid products, including Duragesic (a transdermal fentanyl patch) and Nucynta (tramadol tablets and oral solution). Through its Noramco and Tasmanian Alkaloids subsidiaries, Johnson & Johnson farmed the poppy plant in New Zealand and created the precursor chemical and raw materials necessary to manufacture *all* opioids. Noramco and Tasmanian Alkaloids sold these raw materials to the other opioid manufacturers: Purdue, Endo, Mallinckrodt, and others. Johnson & Johnson was the origin point in the entire opioids supply chain.

408. Just like McKinsey's relationships with Purdue, Endo, and the others, McKinsey's opioid-related work for Johnson & Johnson spanned decades.

409. Just as Endo was led by former partner McKinsey partner Rajiv de Silva, Johnson & Johnson similarly relied on McKinsey as a pipeline for its own management timber. As described above, McKinsey alumni tend to move on to positions with McKinsey clients. Janssen's current

²⁷² MCK-MDL2996-0358871, at 0358872; ENDO-OPIOID_MDL-02201117.

Director of Customer Marketing & Value Based Care was hired from McKinsey's PMP group. The relationship flows both ways: Janssen's former Vice President of Sales and Marketing for Janssen Pharmaceuticals is currently a McKinsey partner. Moreover, Ian Davis has been an independent director since 2010 and currently sits on the Audit and Regulatory Compliance committees of Johnson & Johnson's board. Previously, he was a Senior Partner at McKinsey, "having served as Chairman and Worldwide Managing Director from 2003 until 2009."²⁷³

410. Kevin Sneader, until recently McKinsey's global managing partner, and one of Davis' successors, described Davis as a "mentor" who was the managing partner of McKinsey's London office when Sneader was working there and "worked on one of his teams." 274 Given Frazier's presence on the board, Johnson & Johnson was obviously an important account for McKinsey. At present, it is not known which McKinsey partner(s) was the Director(s) of Client Services for the Johnson & Johnson account.

411. What is known, however, is that McKinsey

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²⁷⁵ On July 6, 2011, Ghatak

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²⁷³ https://www.jnj.com/leadership/ian-e-l-davis

attended an internal McKinsey call with the consultants working on the Johnson & Johnson account

to discuss the "J&J Nucynta sales force disruption." The same day, Laura Moran, who like

Ghatak worked both the Purdue and Endo accounts, also provided internal advice regarding

Nucynta to her McKinsey partner Gerti Pellumbi, who was leading Nucynta sales efforts for the

Johnson & Johnson account, and engagement manager Bryan Reinholt, who was with Pellumbi on

²⁷⁵ JAN-NH-00167575.

²⁷⁴ See Interview with Kevin Sneader, Harvard Project for Asian & International Relations, January 31, 2021, available at https://www.youtube.com/watch?v=qed53EGG8kU

²⁷⁶ MCK-MDL2996-0222833.

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²⁸² MCK-MDL-2996-0261694.

the Johnson & Johnson account. 277 Martin Elling, one of the lead McKinsey partners on the Purdue account alongside Ghatak, attended internal McKinsey calls on March 25, 2010, 278 and again on May 27, 2011 to discuss McKinsey's work for Johnson & Johnson's Nucynta.²⁷⁹ Then, on December 13, 2011, Elling attended a meeting with Johnson & Johnson personnel regarding "acceleration opportunities." 280 Aamir Malik attended the meeting with Elling, and, naturally, also worked on the Endo account.²⁸¹ Malik and Ghatak had an internal McKinsey meeting amongst themselves regarding the "Nucynta Kickoff" at Johnson & Johnson six months prior, on June 3, $2011.^{282}$

1. Noramco

412. Janssen was not the only Johnson & Johnson unit that

and Janssen was not Johnson & Johnson's only division involved in the narcotics

- 413. Opioids—all of them—are derivatives of opium, which is derived from the poppy plant. In order to sell opioids, someone needs to farm the opium poppy and process the harvest into the raw materials necessary for opioid manufacturers—all of them—to make their products.
- 414. Johnson & Johnson was that farmer. It owned Noramco and Tasmanian Alkaloids, which grew poppies in New Zealand and sold the raw ingredients for opioids to practically all manufacturers.

²⁷⁷ MCK-MDL2996-0419348.

²⁷⁸ MCK-MDL-2996-0256186.

²⁷⁹ MCK-MDL-2996-0255907.

²⁸⁰ MCK-MDL-2996-0255926.

²⁸¹ *Id.*; see also MCK-MDL-2996-0348536 (Example of Malik's work on Endo account).

1 On August 19, 2009, McKinsey's 415. 2 3 4 416. 5 6 7 8 9 417. Seven years later, in 2016, Johnson & Johnson exited the business by selling 10 Noramco and Tasmanian Alkaloids to SK Capital, a private equity firm focused on the 11 pharmaceuticals business, for approximately \$800 million.²⁸⁵ 12 13 Not only was McKinsey's advice 14 invaluable to Johnson & Johnson, the perspective McKinsey gained of the overall opioid market 15 16 from advising the principal upstream supplier to the entire industry would be invaluable in to its 17 own work with its other opioid manufacturer clients. 18 2. **Duragesic** 19 Fentanyl was first synthesized by Paul Janssen and his pharmaceutical company 418. 20 Janssen Pharmaceuticals in 1959. In the 1990s, the company (by then owned by Johnson & 21 Johnson) developed Duragesic, which is a transdermal patch that administers fentanyl to the patient 22 wearing it. 23 24 25 26 ²⁸³ NORAMCO_TX_01136410. 27 ²⁸⁴ NORAMCO TX 01136411, slide 4. ²⁸⁵ Gareth Macdonald, "US Investor buys J&J's opiate API business and announces restructuring," Outsourcing 28 Pharma, July 20, 2016, available at: https://www.outsourcing-pharma.com/Article/2016/07/21/US-investor-buys-J-J-

s-opiate-API-business-and-announces-restructuring

419. "Duragesic proved to be one of the most successful analysis pharmaceutical products ever developed, with sales in 2004 (its last year of patent life) exceeding \$2.4 billion. The success of the fentanyl patch caused many generic companies to produce equivalents once it went off patent." ²⁸⁶

- 420. McKinsey was an integral part of fentanyl's success. As early as 2002, McKinsey was advising Johnson & Johnson regarding methods to boost sales of its opioids. For example, on March 14, 2002, McKinsey prepared a confidential report for Johnson & Johnson's subsidiary Janssen regarding how to market their opioid Duragesic. Incredibly, one of the recommendations McKinsey provided to Johnson & Johnson was that they concentrate their sales and marketing efforts on doctors that were *already* prescribing large amounts of Purdue's OxyContin.²⁸⁷
- 421. In other words, as early as 2002, McKinsey had such intricate knowledge of the sales and marketing practices of opioid manufacturers, generally, and Purdue's efforts with OxyContin, specifically, that it was able to recommend to *a competitor of Purdue* that it boost its own opioid sales by *following in the footsteps of Purdue*.
- 422. McKinsey also advised Johnson & Johnson to target Duragesic on "high abuse-risk patients (e.g., males under 40)." This targeting would take advantage of the marketing claim that Duragesic "was harder to abuse than other opioids on the market."²⁸⁸
- 423. McKinsey helped Janssen target its opioid marketing by identifying "priority growth opportunities" and growth strategies for Duragesic.²⁸⁹ In 2002, McKinsey considered "[w]hat are

²⁸⁶ Theodore Stanley, "The Fentanyl Story," The Journal of Pain, Vol. 15, No. 12 (December), 2014, pg. 1220, available at: https://www.jpain.org/article/S1526-5900(14)00905-5/pdf

²⁸⁷ Chris McGreal, *Johnson & Johnson faces multibillion opioids lawsuit that could upend big pharma*, The Guardian, June 23, 2019, *available at:* https://www.theguardian.com/us-news/2019/jun/22/johnson-and-johnson-opioids-crisis-lawsuit-latest-trial

²⁸⁸ Julia Lurie, "Inside Johnson and Johnson's Quiet Domination of the Opioid Market," June 11, 2019, Mother Jones, available at https://www.motherjones.com/politics/2019/06/johnson-and-johnson-opioid-poppies-tasmania-oklahoma-lawsuit/

²⁸⁹ Oklahoma v. Johnson & Johnson, Oklahoma Proposed Findings & Conclusions, citing 5/30/19pm Tr. & S-1253; see also JAN-MS-00481545 (Deem-Eshleman Ex 73).

example;²⁹⁰

²⁹⁰ Oklahoma v. Johnson & Johnson, 5/30/19 Tr. At 46:1-15.

²⁹¹ JAN-MS-00481547 (Deem-Eshleman Ex 74).

²⁹² MCK-MDL-2996-0135636.

3. Turbocharging Nucynta

424. McKinsey's infamous Project Turbocharge to boost OxyContin sales at Purdue in 2013 and 2014—the same project detailed in Purdue's 2020 guilty plea with the Department of Justice—was not McKinsey's first experience turbocharging opioid sales. Before OxyContin, there was Nucynta:²⁹²

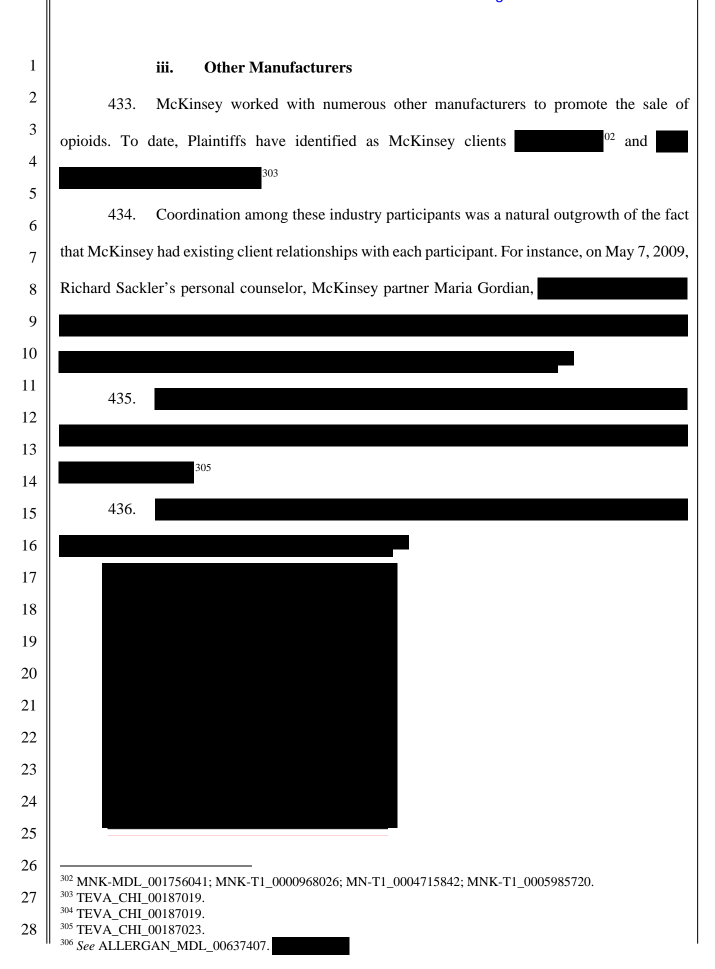
settings of care for opioid high-prescribers and treaters of back pain," listing the "elderly" as an



425. Nucynta was Janssen's branded tapentadol product. Tapentadol is generally regarded as a moderately strong opioid. Nucynta was first approved as a Schedule II controlled opioid agonist tablet and oral solution in 2008 and indicated for "relief of moderate to severe acute pain in patients 18 years of age or older." In 2011, Janssen obtained approval for a long-acting

1 version Nucynta ER, which was indicated for "management of moderate to severe chronic pain in 2 adults and neuropathic pain associated with diabetic peripheral neuropathy (DPN) in adults." 3 426. McKinsey is a repeat opioid sales turbocharger. McKinsey's efforts to turbocharge 4 Nucynta sales resembled those it later deployed in more robust form at Purdue a few years later. 5 For example, "physician prescribing habits," and "switching behavior," were external factors 6 McKinsey identified as key issues "impacting future Nucynta growth." Understanding these issues 7 8 at a granular level would be crucial, including "What is physician/market awareness of Nucynta 9 ER? By physician segment?"²⁹³ These same factors drove McKinsey's later work turbocharging 10 OxyContin. 11 427. Along the way, 12 13 14 15 428. 16 Despite this ambivalence about tamper-resistance, in a status update on June 23, 17 2011, McKinsey informed Janssen that its "initial physician interview findings" indicate Nucynta 18 ER has "lower addictive/abuse potential and side-effect profile as key differentiators vs. Oxycontin 19 ER."295 20 429. As part of the turbocharge process, 21 22 23 24 25 26 27 ²⁹³ MCK-MDL-2996-0135636. 28 ²⁹⁴ JAN-MS-00322271. ²⁹⁵ MCK-MDL-2996-0009526, at 0009529.

³⁰¹ DEPO-CDI-00071072 (emphasis added).



1 iv. McKinsey's Work with Opioid Distributors 2 437. McKinsey worked with opioid distributor 3 4 438. McKinsey worked with opioid distributor McKesson on the company's 5 6 McKinsey's Work with the FDA v. 7 439. As described above, McKinsey assisted Purdue and others to confront FDA 8 9 regulations that posed threats to their clients' ability to maximize revenues from their opioid 10 products. McKinsey's role in shepherding its clients through regulatory interactions takes on a 11 different hew when considered in light of one of McKinsey's other clients: the Food and Drug 12 Administration itself. 13 440. Indeed, the FDA has proved a massive client for McKinsey, who since 2000 has 14 endeavored to expand its public sector practice under the direction and leadership of Nancy 15 Killefer, a now-retired senior partner and director of the firm.³⁰⁹ Since 2008, the FDA has paid 16 17 McKinsey more than \$140 million. 310 A significant portion of that work for the FDA related to the 18 FDA's Center for Drug Evaluation and Research ("CDER"). The CDER is the principal division 19 tasked with approving, among other classes of drugs, opioids. Since 2008, McKinsey has been 20 awarded at least 17 contracts worth at least \$48 million for CDER work.³¹¹ 21 441. The REMS protocols, discussed above, that McKinsey assisted Purdue and others 22 in surmounting beginning in 2008 and culminating in 2012, were overseen by CDER.³¹² 23 24 25 ³⁰⁷ ABDCMDL12135609, slide 5.

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³⁰⁸ MCKSTCT00753097; MCKSTCT00753098.

³⁰⁹ Duff McDonald, *The Firm*. Killefer is also a director of Cardinal Health, one of the distributor defendants in the ongoing nationwide opioid litigation, and a company subject to FDA regulations.

³¹⁰ Letter to Dr. Janet Woodcock from Senator Margaret Hassan et al, August 23, 2021, available at: https://www.hassan.senate.gov/imo/media/doc/fda-mckinsey_letter-final-210823.pdf ("Hassan Letter") 311 Id.

^{28 | 311} Id

³¹² *Id*.

442. Meanwhile, in 2010, McKinsey advised the FDA on building a monitoring system called "track and trace" to assist in the identification of potentially improper distribution of harmful prescription drugs, such as opioids. "The 'track and trace' system deeply impacted McKinsey clients, including the nation's three largest drug distributors—McKesson, AmerisourceBergen, and Cardinal Health [where Killefer has been a director since 2015]."³¹³

- 443. Under one contract, McKinsey developed a roadmap and implemented plans to modernize CDER's new drug regulatory program. Under another, McKinsey developed a framework to increase information technology project delivery across CDER.³¹⁴
- 444. In 2007, Congress passed the Food and Drug Administration Amendments Act ("FDAAA"), which placed new restrictions on the use of certain high risk prescription drugs, including opioids. The new law mandated that FDA require manufacturers of certain drugs to create REMS.
- 445. The FDAAA also required the Secretary of Health and Human Services "to develop standards and identify and validate effective technologies for the purpose of securing the drug supply chain against counterfeit, diverted, subpotent, substandard, adulterated, misbranded, or expired drugs." 21 U.S.C. § 355e(a).
- 446. In 2010 and 2011, under the FDAAA, the FDA awarded McKinsey contracts to design a "track and trace" system to monitor prescription drugs, including opioids, throughout the supply chain and to streamline the drug approval process. The track and trace system had the greatest effect on drug distributors, including McKinsey clients McKesson, AmerisourceBergen, and Cardinal Health.³¹⁵

³¹³ See http://cg.cardinalhealth.com/board-of-directors/default.aspx; Hassan Letter.

³¹⁴ Letter to Senator Chuck Grassley from Andrew Tantillo, Oct. 22, 2021, *available at*: https://www.grassley.senate.gov/imo/media/doc/fda_to_grassley_-_mckinsey_conflicts_of_interest.pdf ³¹⁵ Hassan Letter.

447. Under these contracts, McKinsey was required to consult with "supply chain stakeholders," which likely included these three McKinsey clients as well as pharmaceutical manufacturers.³¹⁶

- 448. In 2011, McKinsey also won a \$1.8 million contract with CDER's Office of Surveillance and Epidemiology ("OSE"), which monitors and evaluates the safety profiles of drugs available to American consumers.³¹⁷ OSE "evaluates more than 2 million adverse event reports submitted every year to FDA's MedWatch program" and provides "risk management expertise on development and implementation of programs and initiatives to support [CDER's] policies related to [REMS] authorities.³¹⁸
- 449. The OSE contract tasked McKinsey with a widespread mission of understanding how OSE functions within the context of a broader system of drug safety in CDER and ultimately developing and implementing a new operating model. In other words, McKinsey helped to restructure a key body that has oversight over the opioid supply chain.
- 450. The 2012 Food and Drug Administration Safety and Innovation Act required the FDA to modernize Sentinel, a system meant to monitor the safety of drugs once they are on the market.³¹⁹ According to the FDA, "Sentinel generates *real-world evidence* to support regulatory actions aimed at protecting the public's health," which in turn "inform[s] healthcare provider decision-making for patients."³²⁰
- 451. A 2014 contract with the FDA charged McKinsey with assessing the "strengths, limitations and appropriate use" of Sentinel. Like the track and trace contract, the Sentinel project

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³¹⁶ *Id*.

³¹⁷ https://www.documentcloud.org/documents/21071060-mckinsey-ose-contract

³¹⁸https://www.fda.gov/about-fda/center-drug-evaluation-and-research-cder/cder-office-surveillance-and-epidemiology

³¹⁹ https://www.documentcloud.org/documents/21071047-r_sentinel_assessment_award_contract_sow-redacted-pr

³²⁰https://www.fda.gov/files/about%20fda/published/Sentinel-System-Overview—-Presentation.pdf; https://www.healthaffairs.org/do/10.1377/hpb20150604.936915/full/

required McKinsey to interview "external stakeholders," including "industry organizations" and "drug and device industry leaders." McKinsey also evaluated how the FDA employees used Sentinel to inform regulatory decision making. 322

- 452. McKinsey performed similar work for the FDA as recently as 2019,³²³ when it signed a contract extension with the agency for work relating to the FDA's efforts to modernize the process by which it regulates new drugs.³²⁴
 - 453. The FDA's drug tracking programs have been panned as failures.³²⁵
- 454. A theme was emerging: as new legislation and regulatory systems were enacted that could have hampered the opioid supply chain, McKinsey stepped in as a key consultant for the FDA. Each time, the new system failed to reign in the out-of-control opioid market. While the FDA was not solely responsible for regulating the opioid industry and McKinsey was not wholly responsible for the FDA's inaction, tools like Sentinel and track and trace could have been implemented in a way to provide new information to combat the country's growing opioid crisis.
- 455. At the same time it was consulting for the FDA, McKinsey was working with its opioid industry clients on how skirt the FDA's regulatory systems.

 $[\]frac{1}{321}$ Ian MacDou

 ³²¹ Ian MacDougall, "McKinsey Never Told the FDA It Was Working for Opioid Makers While Also Working for the Agency," *ProPublica* (Oct. 4, 2021), *available at* https://www.propublica.org/article/mckinsey-never-told-the-fda-it-was-working-for-opioid-makers-while-also-working-for-the-agency
 23 Letter to Bob Sternfels from Representative Carolyn B. Maloney, Nov. 5, 2021, *available at*:

Letter to Bob Sternfels from Representative Carolyn B. Maloney, Nov. 5, 2021, *available at*: https://oversight.house.gov/sites/democrats.oversight.house.gov/files/2021-11-05.CBM%20to%20Sternfels-McKinsey%20re%20Document%20and%20Information%20Request%20%28001%29.pdf

³²³ Ian MacDougall, "McKinsey Never Told the FDA It Was Working for Opioid Makers While Also Working for the Agency," *ProPublica* (Oct. 4, 2021), *available at* https://www.propublica.org/article/mckinsey-never-told-the-fda-it-was-working-for-opioid-makers-while-also-working-for-the-agency

Letter to Bob Sternfels from Representative Carolyn B. Maloney, Nov. 5, 2021, *available at*: https://oversight.house.gov/sites/democrats.oversight.house.gov/files/2021-11-05.CBM%20to%20Sternfels-

McKinsey%20re%20Document%20and%20Information%20Request%20%28001%29.pdf

³²⁵ Sabrina Tavernise, "F.D.A. Faulted for Problems With Drug Tracking" *The New York Times*, Jan. 14, 2016, available at https://www.nytimes.com/2016/01/15/health/fda-faulted-for-problems-with-drug-tracking.html; https://www.gao.gov/assets/gao-16-192.pdf

456. For example, McKinsey advised Purdue on how to soften the FDA's proposed REMS and on coordinating with other opioid manufacturers to advocate against strict oversight.³²⁶ The finalized REMS for opioid products was largely devoid of the restrictions that FDA had initially proposed.³²⁷

- 457. McKinsey's work with the FDA was a key factor in why pharmaceutical industry clients tapped McKinsey for FDA-related work. For example, in endorsing McKinsey's proposed strategy of banding together with other opioid manufacturers, Purdue CEO John Stewart suggested that the consultant itself facilitate the pharmaceutical group's approach to FDA. He wrote: "Perhaps a consultant such as McKinsey who did similar work in the industry and FDA on some aspects of clinical trials or a healthcare-related group that would be interested in playing an active role in the program's development and delivery would be a good choice." 328
- 458. McKinsey performed work for the FDA without disclosing its potential conflicts of interest to the FDA in violation of the contracts between the company and the agency.
- 459. The FDA typically includes conflict of interest clauses in its contracts and relies on contractors to assess and report any conflicts. McKinsey's contracts with the FDA related to CDER processes contained such provisions. One contract required McKinsey to "make an immediate and full disclosure, in writing, . . . of any potential or actual organizational conflict of interest or the existence of any facts that may cause a reasonably prudent person to question the contractor's impartiality because of the appearance or existence of bias." 329

³²⁶ Hassan Letter.

³²⁷ Hassan Letter; Maloney Letter.

³²⁸ Purdue Bankruptcy, Doc. 2166-5, at 58-59.

³²⁹ Ian MacDougall, McKinsey Never Told the FDA It Was Working for Opioid Makers While Also Working for the Agency, *ProPublica* (Oct. 4, 2021), available at https://www.propublica.org/article/mckinsey-never-told-the-fda-it-was-working-for-opioid-makers-while-also-working-for-the-agency

- 460. But McKinsey never disclosed its work on behalf of opioid supply clients to the FDA despite having a hand in developing some of the FDA's most important regulatory processes.³³⁰
- 461. Disclosing its conflicts might have turned off the lucrative tap to not only FDA contracts but also to pharmaceutical industry clients, given the clear value such clients placed on McKinsey's work for the FDA.
- 462. McKinsey's manipulation of regulatory requirements—whether to skirt its own contractual requirements or to bend processes that regulate its clients—is nothing new. McKinsey has come under fire from the Office of Inspector General for the General Services Administration for contract procurement violations³³¹ and from the Justice Department related to violation of Chapter 11 bankruptcy rules.³³² Most recently, six senators have begun to investigate the relationship between McKinsey and the FDA³³³ the House Committee on Oversight and Reform is exploring its abusive conduct in connection with the opioid industry.³³⁴
- 463. As one commentator noted, McKinsey's conduct suggests that it "behaves as if it believes the rules should bend to its way of doing things, not the other way around."³³⁵
 - f. McKinsey's Efforts to Increase the Overall Size of the Opioid Market: the Larger the Pie, the Larger the Slice
- 464. McKinsey advised multiple opioid manufacturers regarding how to grow opioid sales. In order to benefit all its clients, McKinsey engaged in efforts to grow the entire opioid

Id.; Letter to Senator Chuck Grassley from Andrew Tantillo, Oct. 22, 2021, *available at*: https://www.grassley.senate.gov/imo/media/doc/fda_to_grassley_-_mckinsey_conflicts_of_interest.pdf

³³¹ Ian MacDougall, How McKinsey Makes Its Own Rules, *ProPublica* (Dec. 14, 2019), available at https://www.propublica.org/article/how-mckinsey-makes-its-own-rules

³³² Mary Williams Walsh and Emily Flitter, McKinsey Faces Criminal Inquiry Over Bankruptcy Case Conduct, *New York Times*, Nov. 8, 2019, *available at* https://www.nytimes.com/2019/11/08/business/mckinsey-criminal-investigation-bankruptcy.html

³³³ Hassan Letter.

³³⁴ Maloney Letter.

³³⁵ Ian MacDougall, How McKinsey Makes Its Own Rules, *ProPublica* (Dec. 14, 2019), available at https://www.propublica.org/article/how-mckinsey-makes-its-own-rules

market, and not only each individual client's share of it. The theory, basically, is that a rising tide lifts all boats.

465. For example, Purdue incentivized its sales staff "to increase not just sales of OxyContin but also generic versions of extended release oxycodone." Typically, one would not wish to encourage the sales of generic competitors that offer a similar product to one's own. If, however, the goal is to position a company so as to look like an attractive acquisition target, the growth of the overall opioid market is just as important as one's own market share: "Whereas pharma salespeople are usually compensated based on their ability to grow sales of a particular medicine, part of the bonus for Purdue's staff was calculated in relation to the size of the overall market."³³⁶ McKinsey designed that plan.³³⁷

466. This notion that the size of a company's market share is not as important as the size of the *overall* market in which it competes is a core insight of McKinsey's granular approach to identifying corporate growth opportunities. Describing their authors' conclusions in *The Granularity of Growth*, McKinsey stated, "One of their most surprising conclusions is that increased market-share is seldom a driver of growth. They contend, instead, that growth is driven

³³⁶ See David Crow, How Purdue's 'one-two' punch fuelled the market for opioids, Financial Times, September 9, 2018, available at: https://www.ft.com/content/8e64ec9c-b133-11e8-8d14-6f049d06439c

Worth noting is that this strategy of increasing overall opioid sales directly benefitted the Sacklers through their ownership of Rhodes Pharma, a generic opioid manufacturer. Naturally, McKinsey worked with the Sacklers in connection with Rhodes as well, including proposing ideas for synergizing Purdue and Rhodes. *See, e.g.*, MCK-MDL2996-0324955; MCK-MDL2996-0285201. Especially worth noting is that this strategy also benefitted McKinsey's other opioid clients as well. As one observer wrote: "They have a huge amount of inside information, which raises serious conflict issues at multiple levels," stated a former consultant, referring to McKinsey's influential role as advisor to multiple participants in a given industry, such as opioid manufacturing. It "puts them in a kind of oligarchic position." Michelle Celarier, *The Story McKinsey Didn't Want Written*, Institutional Investor, July 8, 2019, *available at:* https://www.institutionalinvestor.com/article/b1g5zjdcr97k2y/The-Story-McKinsey-Didn-t-Want-Written.

For example, in an August 15, 2013 presentation to Purdue management entitled "Identifying OxyContin Growth Opportunities," McKinsey noted that "McKinsey's *knowledge of the ways other pharma companies operate* suggests Purdue should reassess the roles of MSL and HECON Groups – and further drive the salesforce to be more responsive to formulary coverage changes." (emphasis added).

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by where a company chooses to compete: which market segments it participates in . . . the key is to focus on granularity, to breakdown big-picture strategy into its smallest relevant components."338

467. In other words, "Purdue's marketing force was indirectly supporting sales of millions of pills marketed by rival companies."339 "It's the equivalent of asking a McDonald's store manager to grow sales of Burger King and KFC," stated a government official with the Department of Health and Human Services.³⁴⁰

f. McKinsey Portrays Itself as Part of a Solution to a Problem It was Integral in Creating.

468. McKinsey's work on the other side of the aisle—helping clients address opioid abuse and addiction—further proves that it was well aware of the risks of OxyContin, and thus the risks of pushing OxyContin sales and high dose sales, and targeting the highest-volume prescribers. McKinsey advised Purdue on "Project Tango," a 2014 plan to enter the addiction drug market.³⁴¹ McKinsey noted the

469. More than assisting specific clients with addressing the crisis itself, McKinsey saw the ongoing opioid crisis as an opportunity to posture itself as contributing more broadly to *society*. McKinsey likes to think of itself as a change agent capable of solving problems that truly matter, and the opioid crisis is one McKinsey realizes matters. Dr. Sarun Charumilind, a McKinsey partner in Philadelphia, "has led the firm's support to clients and *society* to combat the opioid crisis.³⁴³

The granularity of growth, Book Excerpt, McKinsey & Company, March 1, 2008, available at: https://www.mckinsey.com/business-functions/strategy-and-corporate-finance/our-insights/the-granularity-of-growth ³³⁹ See David Crow, How Purdue's 'one-two' punch fueled the market for opioids, Financial Times, September 9, 2018, available at: https://www.ft.com/content/8e64ec9c-b133-11e8-8d14-6f049d06439c

³⁴¹ See David Armstrong, OxyContin Maker Explored Expansion Into "Attractive" Anti-Addiction Market, ProPublica (Jan. 30, 2019), available at https://www.propublica.org/article/oxycontin-purdue-pharma-massachusetts-lawsuit-antiaddiction-market.

³⁴² PPLPC023000714734.

³⁴³ See https://www.mckinsey.com/our-people/sarun-charumilind

- 470. In Detroit, partner Razili Lewis also helps "clients and *society* combat the opioids crisis." She does so by providing "insights, expertise, analytics, and technology."³⁴⁴
- 471. Over in Cleveland, senior partner Tom Latkovic also "helps clients and *society* combat the opioids crisis." ³⁴⁵
- 472. Kana Enomoto, a senior expert in Washington, D.C., is a "national leader in mental health and substance-use policy," who acted as a "content director" on a study to "raise awareness about opioid-use disorders." She also provided strategic guidance to the United States Surgeon General regarding efforts to "combat the opioid epidemic" when she was his Chief of Staff.³⁴⁶
- 473. McKinsey consistently states that it takes its obligations to society seriously. Indeed, the firm has established a center:³⁴⁷

The Center for Societal Benefit through Healthcare was established to build on the long-standing mission of McKinsey's Public & Social Sector and Healthcare Systems & Services Practices to improve healthcare. The Center's work is funded solely by McKinsey; it is not commissioned by any business, government, or other institution. The Center brings a range of capabilities to bear, including McKinsey's healthcare expertise, advanced analytics, functional knowledge, technology assets, network, and investment capacity.

The Center aspires to collaborate with other organizations to drive positive innovation to improve overall health and well-being and reduce healthcare disparities.

- 474. The Center has focused on addressing the impacts of the opioid crisis on society. One of the metrics that McKinsey uses to track the opioid crisis *as a matter of public health* is the "opioid prescribing rate" per 100 people in every county in the United States.³⁴⁸
- 475. As McKinsey's data visualization makes clear, there is an association between areas with higher opioid prescribing rates and higher instances of opioid use disorder.

³⁴⁴ See https://www.mckinsey.com/our-people/razili-lewis

³⁴⁵ See https://www.mckinsey.com/our-people/tom-latkovic

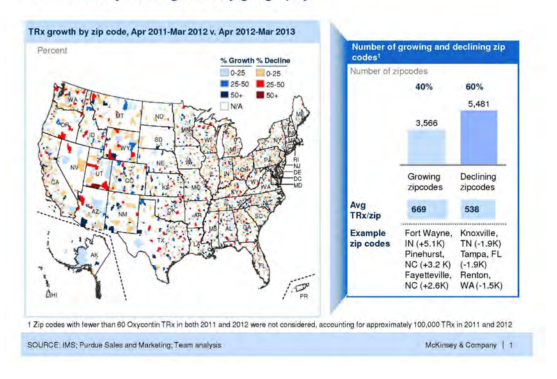
³⁴⁶ See https://www.mckinsey.com/our-people/kana-enomoto

³⁴⁷See https://www.mckinsey.com/industries/healthcare-systems-and-services/how-we-help-clients/center-for-societal-benefit-through-healthcare/overview

³⁴⁸ See https://csbh-dashboard.mckinsey.com/#/datainsights?chart=SC&geo=County&lob=All&metric1=opioid_rxrate&metric2=oud&tab=Map

476. The Center's data visualization is also reminiscent of similar work McKinsey did for Purdue in 2013, although the analysis McKinsey did for Purdue was more granular, analyzing opioid prescribing patterns on the *zip-code* level in all 50 states, as opposed to the county level:³⁴⁹

Exhibit 1: OxyContin growth by geography



³⁴⁹ MCK-MAAG-0024283.

477. In other words, the "opioid prescribing rate" was a metric McKinsey worked with its client to boost for years. Now McKinsey measures the extent of the crisis by the same metric:

Vulnerable Populations: Data By Geography



478. Meanwhile, McKinsey has partnered with Shatterproof, a national non-profit organization dedicated to reversing the addiction crisis in the United States, to prepare a report on overcoming stigma associated with opioid use disorder.³⁵⁰ McKinsey touts the Shatterproof partnership on its webpage as an example of "our societal impact."³⁵¹

479. In August 2017, McKinsey prepared a presentation entitled "Perspectives on Combatting the Opioid Crisis," which referenced its work on combatting opioid addiction for various other entities:

 $^{^{350}\,\}textit{See}\,\, \text{https://www.shatterproof.org/sites/default/files/2020-07/A-Movement-to-End-Addiction-Stigma.pdf}$

³⁵¹ See https://www.mckinsey.com/us/our-societal-impact

RECENT CLIENT EXPERIENCE

- >>> Designed and helped launch a health home program to expand resources and accountability for substance abuse treatment
- Conducted a state wide assessment of opioid prescriber performance in terms of prescribing rate, dosage, and duration
- >> Defined clinically relevant opioid quality measures for a portfolio of episodes-of-care
- Defined clinically relevant opioid quality measures for a Patient Centered Medical Home and Accountable Care Organizations
- >> Used predictive analytics to develop multi-faceted approach to assess patient risk for opioid addiction
- >> Used geo-spatial and social network analytics to assess intensity of opioid abuse and treatment needs
- >> Integrated claims and PDMP data to generate transparency on provider prescribing practices
- >> Developed a substance abuse episode of care focused on priority patient journeys
- 480. In June 2018, Dr. Charumilind and Mr. Latkovic, along with fellow McKinsey partner Elena Mendez-Escobar, published a public report, "Ten insights on the Opioid crisis from claims data analysis," stating information about the risks of opioids that McKinsey knew while advising Purdue to sell more opioids and higher dose opioids, and target the highest volume prescribers:
 - a. "Providers frequently prescribe opioids to patients with known or potential risk factors for abuse[;]"
 - b. "Approximately 35% of the patients given opioid prescriptions in our analysis had features that put them at increased risk for opioid abuse[;]"
 - c. "Most opioids are prescribed by providers other than the natural 'quarterback' of a patient's underlying complaint or condition. . . . This finding makes clear that high-dose prescribers and multi-prescriber patterns are separate issues—and both are important to address[;]" and

- d. "A small portion of opioid use originates in emergency departments." 352
- 481. Two months later, the same authors, joined by Ms. Lewis, published "Why we need bolder action to combat the opioid epidemic." "Our research suggests that much broader and bolder action is required," they announced. 354

h. Coda

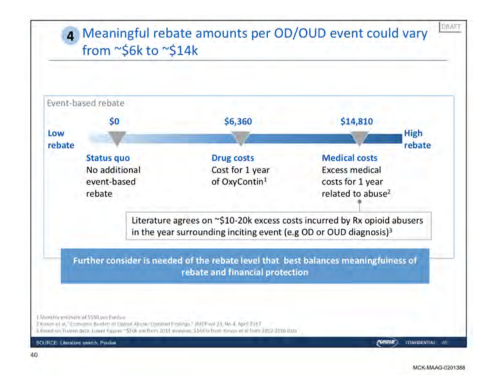
- 482. Marvin Bower, the McKinsey legend who admonished, "Deliver bad news if you must, but deliver it properly," died in 2003, one year before the firm began working with Purdue.
- 483. McKinsey's work with Purdue would have been unrecognizable to Bower, one of the founders of modern management consulting. Instead of acknowledging the elephant in the room—that Purdue's business was knowingly maximizing the amount of addictive and deadly opioids sold in the United States—and delivering that bad news promptly properly to the client, McKinsey instead committed to partner with Purdue to maximize opioid sales without regard to the consequences.
- 484. On October 23, 2017, the president of the United States declared the ongoing nationwide opioid epidemic a "public health emergency." Even at this late hour in the crisis, McKinsey continued to propose solutions to the Sacklers and Purdue to further boost opioid sales. These solutions were fashioned, in perfect McKinsey parlance, as "high impact interventions to rapidly address market access challenges."
- 485. Less than two months after the public health emergency declaration, McKinsey proposed these high impact interventions to Purdue and its board. Among them was perhaps

³⁵² https://www.mckinsey.com/industries/healthcare-systems-and-services/our-insights/ten-insights-on-the-us-opioid-crisis-from-claims-data-analysis

³⁵³ See https://www.mckinsey.com/industries/healthcare-systems-and-services/our-insights/why-we-need-bolder-action-to-combat-the-opioid-epidemic 354 *Id.*

McKinsey's most audacious gambit of the entire Purdue relationship: paying money— "rebates" to health insurers whenever someone overdosed on Purdue's drug.

- 486. These payments for future OxyContin overdoses were christened "Event-Based contracts." 355
- 487. Helpfully, McKinsey provided estimates for the future costs of these "events."³⁵⁶ McKinsey noted that, if Purdue were to start making overdose payments, it would "need to determine which payment amount is optimal."
- 488. A "meaningful" amount, according to McKinsey, would be somewhere between six and fifteen thousand dollars for each person who overdoses or develops opioid-use disorder as a result of Purdue's drugs:



^{355 &}quot;Consultant-ese," when applied to work as grim as maximizing opioid sales in the face of a national disaster, led one former McKinsey consultant to state: "This is the banality of evil, M.B.A. edition." Walt Bogdanich and Michael Forsythe, *McKinsey Proposed Paying Pharmacy Companies Rebates for OxyContin Overdoses*, New York Times, November 27, 2020, *available at*: https://www.nytimes.com/2020/11/27/business/mckinsey-purdue-oxycontin-opioids.html

³⁵⁶ McKinsey defined an "event" as "first occurrence for overdose or opioid use disorder."

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489. The money would be paid to health insurers for the increased costs of additional medical services that resulted from the fact that Purdue's medications caused opioid-use disorder and overdoses in people whose health care costs were the payors' obligation. The money McKinsey proposed Purdue pay out in these circumstances would not go to the individuals afflicted, nor the estates of the dead.

- 490. McKinsey's analysis also suggested that it could predict the number of people who would become addicted to opioids or overdose on pills sold through Purdue's downstream customers. McKinsey "projected that in 2019, for example, 2,484 CVS customers would either have an overdose or develop an opioid use disorder." 357
- 491. It is little surprise, then, that McKinsey was concerned with its legal liability for this work. Within months of recommending "event-based contracts" to Purdue, Martin Elling raised this concern with Arnab Ghatak and suggested corrective action: destroying evidence.

Message

From: Martin Elling [/O=EXCHANGELABS/OU=EXCHANGE ADMINISTRATIVE GROUP

(FYDIBOHF23SPDLT)/CN=RECIPIENTS/CN=6B33C3264F744B04AF05FA59341271BE-MARTIN ELLI]

Sent: 7/4/2018 12:10:13 PM

To: A G [drarnabghatak@gmail.com]

Subject: Re: [EXT]Re: Howdy

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Have a great fourth. M
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> On Jul 4, 2018, at 2:01 PM, A G <drarnabghatak@gmail.com> wrote:

> Thanks for the heads up. Will do.

>> On Jul 4, 2018, at 7:57 AM, Martin Elling <martin_elling@mckinsey.com> wrote:

>> Just saw in the FT that Judy Lewent is being sued by states attorneys general for her role on the Purdue Board. It probably makes sense to have a quick conversation with the risk committee to see if we should be doing anything other that eliminating all our documents and emails. Suspect not but as things get tougher there someone might turn to us. M

>> copy it, disclose its contents or use it for any purpose.

³⁵⁷ Walt Bogdanich and Michael Forsythe, *McKinsey Proposed Paying Pharmacy Companies Rebates for OxyContin Overdoses*, N.Y. Times (Nov. 27, 2020, updated Dec. 17, 2020), https://www.nytimes.com/2020/11/27/business/mckinsey-purdue-oxycontin-opioids.html

492. Elling's prediction that things would "get tougher" for Purdue would prove prescient.

i. Guilty Again - 2020

- 493. On October 20, 2020, Purdue—McKinsey's co-conspirator—agreed with the United States Department of Justice to plead guilty to improper marketing of OxyContin and other opioids again (the "2020 Settlement Agreement"). This time the plea agreement concerned conduct from 2010 to 2018. The agreement includes \$8.3 billion in penalties from Purdue and \$225 million from the Sackler family.
- 494. Purdue pleaded guilty to a dual-object conspiracy to defraud the United States and to violate the Food, Drug, and Cosmetic Act, 21 U.S.C. §§ 331, 353, violating anti-kickback laws, and "using aggressive marketing tactics to convince doctors to unnecessarily prescribe opioids—frivolous prescriptions that experts say helped fuel a drug addiction crisis that has ravaged America for decades."
- 495. The new plea agreement does not identify Purdue's co-conspirators, and McKinsey is not identified by name in the agreement. Instead, McKinsey is referred to as the "consulting company."
- 496. Purdue's new guilty plea concerns Covered Conduct (as defined in the plea agreement) that directly implicates McKinsey in the conspiracy. It is the same conduct described in this Complaint.
- 497. Indeed, the plea agreement signed by McKinsey's co-conspirator states bluntly: "Purdue, *in collaboration with [McKinsey]*, implemented many of [McKinsey's] recommendations." (emphasis added).

498. Further, Purdue admitted that E2E "was overseen by [McKinsey] and some of Purdue's top executives through the creation of the E2E Executive Oversight Team ('EOT') and Project Management Office ('PMO')" (emphasis added).

ii. A Mea Culpa

499. On December 5, 2020, six weeks after Purdue's second guilty plea, McKinsey issued a rare public statement regarding its work with a specific client on its website. The client was Purdue, and the statement was issued is response to Purdue's second guilty plea and recent media reports regarding McKinsey's work selling OxyContin after 2007:

McKinsey statement on its past work with Purdue Pharma

December 5, 2020—As we look back at our client service during the opioid crisis, we recognize that we did not adequately acknowledge the epidemic unfolding in our communities or the terrible impact of opioid misuse and addiction on millions of families across the country. That is why last year we stopped doing any work on opioid-specific business, anywhere in the world.

Our work with Purdue was designed to support the legal prescription and use of opioids for patients with legitimate medical needs, and any suggestion that our work sought to increase overdoses or misuse and worsen a public health crisis is wrong. That said, we recognize that we have a responsibility to take into account the broader context and implications of the work that we do. Our work for Purdue fell short of that standard.

We have been undertaking a full review of the work in question, including into the 2018 email exchange which referenced potential deletion of documents. We continue to cooperate fully with the authorities investigating these matters.

- 500. As the statement indicates, McKinsey stopped doing work "anywhere in the world." Given that Purdue's operations addressed only the United States, the global reach of McKinsey's regret is noteworthy.
- 501. In August 2013, when the Sacklers adopted McKinsey's "Project Turbocharge" for Purdue, Tim Reiner, a long-time McKinsey consultant, joined Mundipharma. Mundipharma is a separate company—also owned by the Sacklers—that sells opioids internationally.

As late as 2019, Mundipharma has been asserting many of the same misleading

claims about opioids that previously led to criminal liability in the United States.³⁵⁸ McKinsey has

long assisted the Sacklers in growing Mundipharma's opioids market.³⁵⁹ By 2015, McKinsey's

workload with Mundipharma was large enough to merit formal coordination and incorporation with

the overall McKinsey team servicing the Purdue account. Around this time, McKinsey's Elling

agreed to assume "a real operational DCS" role with respect to the work that McKinsey was

performing for the various Sackler interests, including "integrat[ing] the Mundipharma stuff." ³⁶⁰

Even if the various components of the Sackler "family conglomerate" were nominally independent,

McKinsey consolidated its own treatment of its work for all of these companies as serving just a

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single client.

iii. A Hedge Fund

502 On Echman 4 2021

503. On February 4, 2021, forty-nine state attorneys general announced a multistate settlement with McKinsey related to its work for opioid manufacturers. McKinsey agreed to pay almost \$600 million dollars. At the time of the announcement, most of the participating states each filed a complaint and consent decree finalizing the settlement.

504. Three days after the settlement, it came to light that McKinsey appears to have benefitted from its work promoting opioids not only through the fees paid to McKinsey by its clients, but also through investments in opioid-related business made by McKinsey's own hedge fund, the McKinsey Investment Office ("MIO"). MIO is the hedge fund referenced above, with respect to McKinsey's investment in Teva Pharmaceutical.

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³⁵⁸ See Kinetz, Erika, Fake doctors, pilfered medical records drive OxyChina sales, Associated Press, November 19, 2019, available at: https://apnews.com/article/4122af46fdba42119ae3db30aa13537c

³⁵⁹ See, e.g., MCK-MDL2996-0256120; MCK-MDL2996-0327127; MCK-MDL2996-0183279; MCK-MDL2996-0238998; MCK-MDL2996-0286490.

³⁶⁰ MCK-MDL2996-0210149

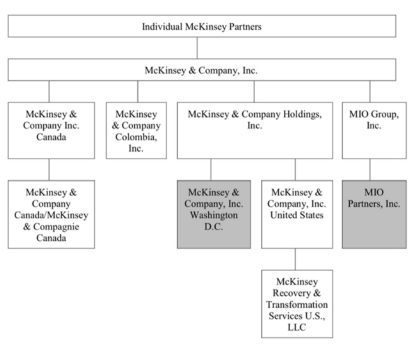
505. Consultants don't typically have in-house hedge funds overseeing retirement accounts and partners' personal investments. In fact, McKinsey is the only one. "Most large companies, including all the major consulting firms, hire third-party firms . . . to oversee their employees' retirement accounts." MIO manages approximately \$31 billion on behalf of McKinsey partners, employees, and former partners.

- 506. Through MIO, McKinsey was heavily invested in the opioid industry, and stood to gain financially from the continuation of the opioid crisis. It even invested in opioid addiction treatment businesses—a growing industry, as McKinsey knew.
- 507. In short, "during the years McKinsey was helping opioid makers propel sales of the drugs, MIO Partners held stakes in companies that profited from increased usage." ³⁶³
 - 508. To understand MIO, an organizational chart of McKinsey is helpful:

³⁶¹ Gretchen Morgenson, "Consulting giant McKinsey allegedly fed the opioid crisis. Now an affiliate may profit from treating addicts.," *NBC News*, February 8, 2021, *available at* https://www.nbcnews.com/news/us-news/consulting-giant-mckinsey-allegedly-fed-opioid-crisis-now-affiliate-may-n1256969

SEC Order dated November 19, 20201 at Para. 5, available at: https://www.sec.gov/litigation/admin/2021/ia-5912.pdf. That \$31 billion under management would make MIO Partners the thirteenth largest hedge fund on Earth. See https://www.pionline.com/interactive/largest-hedge-fund-managers-2021.

³⁶³ Gretchen Morgenson, "Consulting giant McKinsey allegedly fed the opioid crisis. Now an affiliate may profit from treating addicts.," *NBC News*, February 8, 2021, *available at* https://www.nbcnews.com/news/us-news/consulting-giant-mckinsey-allegedly-fed-opioid-crisis-now-affiliate-may-n1256969



- 509. MIO Group, Inc., and MIO Partners, Inc. are directly-owned subsidiaries of McKinsey & Company, Inc. Given that McKinsey advises countless large corporations, McKinsey's hedge fund inevitably invests in McKinsey's clients.
- 510. MIO manages money for pension plans sponsored by McKinsey in which current and former McKinsey employees participate, as well as privately-offered investment funds available to partners and former partners. Today, nine of MIO's eleven directors are current or former McKinsey partners. Prior to 2017, there were no outside directors at MIO.
- 511. MIO structures its investment activities in three principal ways: (1) approximately 50-60% of MIO's assets are managed by third-party money managers, who have sole discretion on what securities to buy with MIO's money, and where MIO may or may not have information regarding which securities the third-party money manager has purchased for MIO's benefit; (2) "separately managed accounts," comprising approximately 40% of MIO's holdings, are portfolios of securities managed by a third-party money manager, but where MIO "knows what securities are held through each account," and; (3) direct investments, where MIO invests its own money directly, which comprises approximately 10% of MIO's investments.

512. In other words, for *at least 40*% of MIOs holdings, McKinsey partners are able to know the specific investments held by the various MIO funds. "MIO has a ledger for every security in their managed accounts."³⁶⁴ That comprises a pool of capital worth more than \$6 billion.

513. MIO is run for the benefit of McKinsey's partners and, to a separate extent, McKinsey's employees. Those individuals (and, crucially, former McKinsey partners) invest their own money in MIO, and their access to those investment opportunities constitutes a meaningful and important component of those individuals' compensation. MIO has, "at a minimum, the ability to view the individual securities that account for approximately 40 to 50 percent." This is approximately \$6 billion dollars of invested capital. What is more, MIO *directly invests* approximately 10% of its assets. That is \$1.5 billion MIO directly invests in securities without the use of any outside money manager. These numbers exclude leverage.

- 514. From a conflicts perspective, the fact that *former* partners may participate in MIO investments merits consideration. With respect to McKinsey's opioid investments, it is notable to consider just who some of those "former partners" are. As noted above, Rajiv de Silva, Chief Executive Officer of opioid defendant Endo Pharmaceuticals, is a former McKinsey partner. Kare Shultz, Chief Executive Officer of opioid defendant Teva Pharmaceutical, is a former McKinsey partner. Frank Scholz, President of opioid defendant SpecGX, a subsidiary of Mallinckrodt, is a former McKinsey partner. Marc Owen, President of opioid defendant McKesson, is a former McKinsey partner. This list is merely illustrative; it is not exhaustive.
- 515. The result is the prospect of individual executives at various opioid manufacturing and distribution companies obtaining financial gain from the ongoing propagation of the opioid

Michelle Celarier, *McKinsey's Managed Accounts Come Under Scrutiny in Trial*, Institutional Investor, February 5, 2020, *available at:* https://www.institutionalinvestor.com/article/b1k6wnn251s472/McKinsey-s-Managed-Accounts-Come-Under-Scrutiny-in-Trial

crisis *not* via compensation from their employers, but via participating in investments alongside their *former* employer (and, in many cases, current consultant).

- 516. Three days after McKinsey and the state attorneys general announced their settlement, NBC News reported that MIO, McKinsey's hedge fund, owned opioid-related investments during the time that it advised its opioid clients.
- 517. One is Deerfield Management Co., "a \$10 billion dollar health care investment firm based in New York." As ever, "two top Deerfield executives previously worked at McKinsey." A retirement fund managed by MIO held a \$108 million stake in funds managed by Deerfield and invested in opioid industry participants. "In 2017, for example, Deerfield was a 6 percent shareholder in Mallinckrodt, a major opioid maker." From 2011 through 2016, Deerfield held a stake of up to \$90 million in Teva. Deerfield also took stakes in the distributors described above, including McKesson and Cardinal Health. 367
- 518. McKinsey is also invested in treatment, an inevitable growth industry sprouting from the over-selling of opioids. Separate from its investments with Deerfield, MIO is also invested in Adamis Pharmaceuticals, "a company that develops products to treat opioid overdoses," and therefore "may also benefit from opioid settlement funds" paid by McKinsey as a result of its settlement with the states. As of 2020, MIO owned 26% of the Adamis' preferred shares through another outside investment manager (not Deerfield). Separately, Deerfield invested \$331 million in Recovery Centers of America, an addiction treatment company that operates facilities in states that McKinsey recently settled with.

³⁶⁵ Gretchen Morgenson, "Consulting giant McKinsey allegedly fed the opioid crisis. Now an affiliate may profit from treating addicts.," *NBC News*, February 8, 2021, *available at* https://www.nbcnews.com/news/us-news/consulting-giant-mckinsey-allegedly-fed-opioid-crisis-now-affiliate-may-n1256969

 $^{27 \}parallel ^{366} Id.$

³⁶⁷ *Id*.

³⁶⁸ *Id*.

³⁶⁹ *Id*.

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519. These relationships and investments give a glimpse into the myriad means McKinsey deploys to make money. Consulting is more than giving advice. Indeed, On November 19, 2021, MIO Partners agreed to pay an \$18 million fine to the SEC due to MIO's possession of material nonpublic information related to its holdings, information obtained through consulting.

i. **Publicis: "The Power of One"**

- 520. Defendant Publicis Health is an advertising and consulting company that services pharmaceutical manufacturers. Publicis Health is a division of the French multi-national advertising and communications conglomerate Publicis Groupe, S.A. ("Publicis"). Annual revenues exceed \$9 billion annually.
- 521. Publicis is one of the "Big Four," as the four firms that account for more than half of the global advertising industry are called.³⁷⁰ In 2002, as the opioid crisis was taking root across the United States, the president of the American Association of Advertising Agencies, stated, "Now you have four megacompanies with revenues that are staggering, bigger than some of the companies they serve."371
- 522. The rise of the Big Four came through decades of mergers and acquisitions of separate agencies and industry consolidation; each is essentially a conglomerate. For its part,

³⁷⁰ The other three are Omnicom Group, the Interpublic Group of Companies, and the WPP Group. Stuart Elliott, Advertising's Big Four: It's Their World Now, New York Times, March 31, 2002, available at: https://www.nytimes.com/2002/03/31/business/advertising-s-big-four-it-s-their-world-now.html

Other primary vendors to large companies have their own little linguistic conferences. For advertising's Big Four, consulting has its Big Three - also referred to as "MBB" - comprised of McKinsey, Bain, and Boston Consulting Group. See https://en.wikipedia.org/wiki/Big Three (management consultancies). The Big Four and the Big Three increasingly compete for the same project work at the same clients. Moreover, accountancies have their own Big Four, and compete with the Big Three and the other Big Four. See https://en.wikipedia.org/wiki/Big Four accounting firms. These Big vendors provide services demanded by thousands of overlapping clients. That market reality confers substantial power to this Big, elite few.

³⁷¹ Stuart Elliott, Advertising's Big Four: It's Their World Now, New York Times, March 31, 2002, available at: https://www.nytimes.com/2002/03/31/business/advertising-s-big-four-it-s-their-world-now.html

Publicis grew by acquiring the agencies Fallon McElligott, Saatchi & Saatchi, and Nelson Communications, among many others.³⁷²

- 523. Today, Publicis operates numerous subsidiaries focusing on subsets of the advertising industry. Publicis Health is the conglomerate's division that specializes in work for healthcare and pharmaceutical companies. Within Publicis Health, there at least fifteen "agency brands" subsidiaries, essentially, operating under their own brand names. Each provides specialized advertising and communications strategies to Publicis' pharmaceutical clients. Razorfish Health, Publicis Health Media, Digitas Health, Rosetta, and Verilogue are some examples.³⁷³
- 524. Each brand specializes in a specific niche within the overall suite of sales and marketing services offered by Publicis Health. Razorfish pioneered and specializes in digital marketing; Digitas specializes in interactive marketing; Publicis Health Media's wheelhouse has been creative and media marketing. Verilogue's niche is providing audio recordings of interactions between patients and doctors that may be mined for insights on how to sell more drugs. Routinely, a Publicis client would engage with more than one of its subsidiaries in tandem and as part of an overall client relationship with Publicis.³⁷⁴
- 525. Until 2019, Publicis also owned Publicis Touchpoint Solutions, which provided contract sales organization ("CSO") services to pharmaceutical manufacturer clients.³⁷⁵ Pharmaceutical companies routinely seek to optimize their salesforces to maximize profitability.

³⁷² Stuart Elliott, Advertising's Big Four: It's Their World Now, *New York Times*, March 31, 2002, *available at*: https://www.nytimes.com/2002/03/31/business/advertising-s-big-four-it-s-their-world-now.html
³⁷³ *See* https://publicishealth.com/companies.

³⁷⁴ For example, upon information and belief, Purdue used Publicis Health Media and Razorfish during the same year, and paid separate invoices to each agency, despite their joint ownership.

³⁷⁵ On January 31, 2019, Publicis Healthcare Solutions, formerly known as Publicis Touchpoint Solutions, was sold by Publicis Groupe to Altamont Capital Partners. *See* "Altamont Capital Partners Acquires Publicis Healthcare Solutions," January 31, 2019, *available at*: https://www.prnewswire.com/news-releases/altamont-capital-partners-acquires-publicis-healthcare-

 $[\]underline{solutions 300787750.html\#:\sim:text=Altamont\%20Capital\%20Partners\%20Acquires\%20Publicis\%20Healthcare\%20Solutions}$

As such, the typical pharmaceutical company does not maintain under-utilized salesforces, or salesforces larger than necessary to maximize revenue on the company's current product offerings. The result, oftentimes, is that a pharmaceutical company hoping to launch a new product will not have the resources in-house to adequately push a new product launch.

- 526. Publicis Touchpoint Solutions solved these problems for clients by offering contract salesforces to augment the number of sales representatives a manufacturer can deploy in order to maximize the success of a product launch. Or, in many cases, Publicis would employ and control the entire sales force for a given drug, on a contract basis, for drug manufacturers that wish to outsource the entirety of their sales and marketing efforts.³⁷⁶ As will be seen, Publicis Touchpoint Solutions provided sales representatives to numerous opioid manufacturers for numerous opioid products at different stages of the product life cycle.
- 527. These different divisions offering complementary services to clients function as a seamless whole. "The Power of One' is Publicis Groupe's operating philosophy. Bringing together 80,000 employees across more than 110 countries and 56 agency brands, we deliver a seamless and modular experience in the relentless service of our clients," Publicis says.³⁷⁷ That seamless and modular experience is "free from silos," with "unified P&L's" and no operational barriers between Publicis' brands.³⁷⁸
- 528. "The Power of One" is more than marketing pablum. It governs the operations of the parent organization and how it exerts control over its numerous agencies. This control can at times lack subtlety. "Publicis Groupe executives gathered a few months ago to debate which agency would service a new piece of business won by the holding company's centralized Power of One

³⁷⁶ While the new product launch is a classic use case for a CSO, is it's not their only use. A drug manufacturer may also choose to utilize a CSO at other stages of the product life cycle, for instance as loss of exclusivity approaches and a manufacturer wishes to re-deploy its internal sales force to focus on newer drugs or those about to be launched.

³⁷⁷ See https://publicishealth.com/companies.

³⁷⁸ See https://www.publicisgroupe.com/en/the-groupe/about-publicis-groupe

team, which is composed of talent from its various shops. According to a person at the meeting – a former creative from a Publicis agency – a suggestion was made for the assignment to be handled out of Saatchi & Saatchi New York. According to that creative, one of Publicis' CEO-Chairman Arthur Sadoun's 'main people' responded: 'Don't put that there; [Saatchi] won't be here next year.'"³⁷⁹

529. By October of 2021, Publicis had risen to become the largest advertising conglomerate in the world, with a market capitalization of a little more than \$16 billion.³⁸⁰

j. What Publicis Does: Marketing and Consulting

530. Traditionally, the advertising industry organized itself based on the "agency model," with advertising agencies performing both creative and advertising placement services. At the top of the pyramid was the "agency of record," or "AOR." The AOR is the advertising agency appointed by the client to coordinate the purchase of media time and space for the placement of client advertisements. While any given client may choose to employ multiple advertising agencies to assist with specific projects, the AOR sits atop those other agencies performing project work and directs the placement of project work performed by other agencies for the client. Typically, the AOR will receive payment from an agency performing project work for the client for its placement. Given its control over the placement of client content – in effect, a monopoly on the client marketing distribution channel – the AOR was in a position to influence the conduct of other contracting agencies performing discrete projects for the client, should the client choose to utilize different agencies for different marketing functions (digital vs. print media, for example).

³⁸⁰ See https://www.prweek.com/article/1731568/publicis-overtakes-rivals-worlds-valuable-agency-group

³⁸¹ See https://www.allbusiness.com/barrons_dictionary/dictionary-agency-of-record-4962111-1.html

531. Publicis' various divisions performed multiple roles for various opioid manufacturers, including AOR roles on numerous campaigns as well as project work under the aegis of other AORs. Publicis' Rosetta, for example,

Publicis' Saatchi & Saatchi Healthcare

- 532. The traditional AOR model has evolved significantly over the years as a result of consolidation within the industry, as well as competition from without. As the marketing industry has consolidated into the Big Four, it has transformed its product offerings in response to competition from consulting firms. The result is that Publicis has outgrown its heritage and is now far more than an advertising agency. It has transformed into an offeror of strategy and consulting services in addition to traditional agency work.³⁸³
- 533. In response to competition from outside the traditional agencies, coupled with the expansion of traditional advertising channels to encompass new technologies, the rise of digital marketing, the erosion of traditional print mediums, and other market dynamics, the Big Four agencies have evolved in recent years beyond the traditional AOR hierarchy to offer services to their clients in more nuanced and flexible ways, and to pair traditional product offerings with newer services.

k. A Consulgency

- 534. Publicis is no exception; indeed, it is an exemplar of these industry trends.
- 535. These days, Publicis is not just a content creator and placement agent; it is also a consultant. Indeed, the evolution of management consulting and advertising (and *particularly* pharmaceutical advertising) has created a market reality in which there is substantial overlap in the product offerings of global advertising agencies like Publicis and global management consultants

³⁸² Transcript of Amanda Stephens Hogan Deposition dated January 25, 2019, Pg. 414:3-8,

³⁸³ For example, Publicis Sapient's Strategy & Consulting Practice Group describes its approach, "Our strategy and consulting teams work seamlessly with our experience and engineering teams to ensure we develop the most high-impact strategies to drive effective digital business transformation."

like McKinsey & Company, Inc. Just as McKinsey's traditional management consulting business has evolved to encompass various operational roles performed for clients, including sales and marketing design and implementation, the traditional advertisers have similarly evolved.

536. Publicis, for example, evolved to offer strategy consulting services and perform implementation work in addition to merely proposing advertising campaigns. By 2018, Publicis' investor presentation emphasized this shift, declaring that "We have the organization to shift from a communications partner to a transformation partner." The goal of any Publicis client relationship is to "be our clients' indispensable partner in their transformation." 385

537. This is part of a broader shift in the overall consulting and marketing industries. "Traditionally, marketing agencies spoke to the Chief Marketing Officer and implemented communication strategy, while consultants spoke with the CEO and devised the general strategy – with marketing communications being the tail of it. In an effort to compete, agencies have started to develop consulting skills, creating 'consulgencies.'"386 As an example, Publicis' Publicis Sapient division was ranked in 2019 as the premier leader in providing "digital transformation" services, besting traditional consultants Accenture, Ernst & Young, PwC, and McKinsey.³⁸⁷ According to the rankings analysis, these digital transformation leaders "blend strategy and execution chops and couple them with the soft skills for inspiring leadership and training teams," and identified Publicis as "the top consultancy on the customer experience front" and "an especially strong partner where the transformation emphasis is on creating world-class digital customer and employee experiences."388

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³⁸⁴ **Publicis** Groupe Investor Day Presentation dated March 20. 2018, available at: http://documents.publicisgroupe.com/events/2018-03-investor-day-intro.pdf (emphasis in original).

^{386 &}quot;Agencies, consulting firms, and the future of marketing," Rewire Mag, available at: https://rewire.ie.edu/agencyvs-consulting-firms-future-marketing/

³⁸⁷https://www.consulting.us/news/1991/publicis-sapient-named-a-leader-in-digital-transformation

^{388 &}quot;Publicis Sapient named leader in digital transformation," Consulting.us, March 21, 2019, available at: https://www.consulting.us/news/1991/publicis-sapient-named-a-leader-in-digital-transformation. Firms like Razorfish

- 538. One marketing executive, upon his company being acquired by a large consulting firm, explained, "we see the next decade as belonging to what we call 'consulgencies' offerings at the cusp of what consultancies do and agencies do bringing the best of both worlds together." One benefit of this hybrid model is greater access to client information than a traditional agency model would entail. "I think consultancies are allowed more to look at the business side of things than agencies. Clients are not willing to share confidential information with us," explained one marketing professional. 390
- 539. A natural outcome of this convergence between the management consulting and advertising industries has been partnerships between incumbent management consultants and incumbent agencies as both find themselves competing for the same business in a new competitive landscape.
 - 540. Publicis and McKinsey routinely partner on projects together for mutual clients.
- 541. "McKinsey & Company and Publicis Health work together to help clients develop agile approaches to winning launches," declaimed a pharmaceutical industry marketing publication.³⁹¹ In 2016, Janet Winkler, then a Group President at Publicis Health³⁹², and Brian Fox,

and Sapient were seen as competitors to McKinsey from the outset, as McKinsey was experiencing a wave of employee departures during the early 2000's. "It wasn't just dot-com startups that were alluring. A whole new class of consulting firm burst onto the scene, with hipper names – Razorfish, Scient, Viant, and Sapient – and sexier projects. The work they were doing seemed far more crucial than redrawing organizational charts. They were helping companies use the Internet to transform everything about the way they did business – from sourcing to distribution to how they treated and served their customers." Duff McDonald, *The Firm*, Pg. 265. Razorfish, Scient, and Sapient were eventually acquired by Publicis. (Viant, which was *not* scooped up by one of the Big 4, was instead acquired in 2002 by a company named Divine that liquidated in bankruptcy the following year.)

³⁸⁹ "The deal that unlocks the value of our industry," *Marketing Magazine*, June 8, 2018, *available at*: https://marketingmagazine.com.my/arrival-of-the-consulgency/

³⁹⁰ Shareen Pathak, "'We're giving the business away to consultants:': Agencies brace for new competition, *Digiday*, October 25, 2017, *available at*: https://digiday.com/marketing/giving-business-away-consultants-agencies-brace-new-competition/

³⁹¹ Janet Winkler, Brian Fox, et. al. "Five Inconvenient Truths That Can Make or Break a Product Launch," PM360, November 23, 2016, *available* at: https://www.pm360online.com/five-inconvenient-truths-that-can-make-or-break-a-product-launch/

³⁹² In 2017, Winkler left her role as Group President of Publicis. The following year, she joined McKinsey & Company as a Senior Advisor.

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a Senior Partner at McKinsey, co-authored the article "Five Inconvenient Truths That Can Make or Break a Product Launch" in industry trade publication PM360.

- McKinsev and Publicis co-authored³⁹³ the piece to highlight their joint expertise in 542. pharmaceutical marketing. "Applying complementary tool and capabilities, [McKinsey and Publicis Health] deliver rapid, actionable analytics as well as change management and execution support to accelerate brand performance throughout the lifecycle," explained a note accompanying the article.
- Purdue is one example, but Publicis and McKinsey had many other mutual opioid 543. manufacturer clients. They worked for these clients contemporaneously and for years while the opioid epidemic grew to its present scourge.

l. **Publicis and Purdue: Selling the Sackler Strategy**

As explained in detail above, in the wake of Purdue's 2007 guilty plea with the 544. Department of Justice and accession to a 5-year Corporate Integrity Agreement with the Office of Inspector General for the United States Department of Health and Human Services, Purdue faced newly imposed constraints on its sales and marketing practices. As a result, the Sackler family desired to achieve distance from this "concentration of risk" by diversifying the family fortune away from Purdue, and by increasing OxyContin sales in the near term in order to achieve that distance.

³⁹³ In addition to Winkler and Fox, "Gregory Graves, Associate Partner, McKinsey & Company; Catherine Mayone, EVP, General Manager, Publicis Health and Sapient Health; and Dan Tinkoff, Partner, McKinsey & Company, also contributed to this article." See https://www.pm360online.com/five-inconvenient-truths-that-can-make-or-break-aproduct-launch/

Notably, the McKinsey authors – Fox, Graves, and Tinkoff –

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545. Alongside McKinsey, Publicis was integral to the sales and marketing campaigns deployed to increase OxyContin sales notwithstanding the Corporate Integrity Agreement and in furtherance of the Sacklers wishes.³⁹⁴

i. Hopelessly dependent on consultants; "Desperately seeking new growth."

- 546. Purdue was captured. After the 2007 guilty plea, the Sackler family wished to dispose of Purdue because it was concerned about both legal liability and reputational risk associated with owning the monoline opioid manufacturer. After 2007, the name of the game was profit maximization of a drug Purdue's owners knew had no long-term future.
- 547. The company, in other words, was a basket case: pursuing internally contradictory goals to maximize opioid sales because its previous efforts to maximize opioid sales had created an existential crisis for the company. Threading a needle like that is hard. Purdue needed others to tell it what to do in order to achieve its owners' mandate.
- 548. By the end of 2013, Purdue was seeking any opportunity to maintain growth it could identify. Publicis noted that Purdue's Board (meaning the Sackler family) was "always looming," and the pressure was always to increase sales. One risk Publicis identified was the risk that the Sacklers would fire the CEO at that time, John Stewart.
- 549. Publicis summed up its outlook for its client, "PURDUE in 2014: Desperately seeking new growth."
- 550. This is red meat to consultants. McKinsey & Company. ZS Associates. Publicis. These happy warriors³⁹⁵ serviced Purdue after the company's executives pleaded guilty in 2007,

³⁹⁴ To emphasize the disregard with which the CIA was met, Dr. Richard Sackler has testified, incredibly, that he has never read the agreement, despite his years of continued service on Purdue's Board of Directors before, during, and after the CIA compliance period.

³⁹⁵ Consultants love martial metaphors. See "Publicis 2020: Sprint to the Future," Publicis Groupe, March 20, 2018, https://www.publicisgroupe.com/en/news/press-releases/publicis-2020-sprint-to-the-future-en-1 available (Publicis helping clients to "reduce their costs and win the battle with new competition); see also, "A battle plan for telcos' digital attacker brands," McKinsey March 5, 2021, available & Co., at:

and with full knowledge of that guilt.³⁹⁶ All worked for Purdue on an ongoing basis right up until Purdue stopped marketing OxyContin in 2018. And they worked together. They teamed up to service a client so dependent on their offerings that it could no longer function without their ongoing assistance.

- 551. Indeed, the consultants banded together to achieve their own goals with respect to mutual clients. "McKinsey & Company and Publicis Health work together to help clients develop agile approaches to winning launches. Applying *complementary* tools and capabilities, they deliver rapid, actionable analytics as well as change management and execution support to accelerate brand performance."397
- 552. As will be seen, Publicis' complementary tools and capabilities were critical to McKinsey's *Project Turbocharge*, which was adopted by Purdue and implemented by all three (and ZS) in late 2013 and thereafter.

ii. **Branded Marketing**

553. Purdue was a monoline manufacturer of opioids. OxyContin (oxycodone), Hysinglia (hydrocodone), Targiniq (oxycodone/naloxone) and Butrans (buprenorphine) were Purdue's principal branded opioid products. Publicis worked on campaigns for all four. But OxyContin was the cash cow. As Publicis' Rosetta unit described it in 2014, "for 17 years Purdue has relied almost solely on the revenue from their \$3b blockbuster opioid medication, OxyContin (~90% of Purdue's revenue)."

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https://www.mckinsey.com/industries/technology-media-and-telecommunications/our-insights/a-battle-plan-fortelcos-digital-attacker-brands;

³⁹⁶ For example, an internal Publicis Client Service Team review document prepared in 2013 for the Purdue account

began with a section entitled, "To know Purdue," which stated, "In 2007, John Brownlee U.S. Attorney charged that 'Purdue, under leadership of its top executives, continued to push a fraudulent marketing campaign... In the process scores died as a result of OxyContin abuse and an even greater number of people became addicted to OxyContin; a drug that Purdue led many to believe was safer, less abusable and less addictive than other pain medications on the market."

³⁹⁷ Janet Winkler and Brian Fox, "Five Inconvenient Truths That Can Make or Break a Product Launch," PM360, November 23, 2016, available at: https://www.pm360online.com/five-inconvenient-truths-that-can-make-or-break-aproduct-launch/ (emphasis added).

554. The Publicis-Purdue relationship began as early as April 2010, during the pendency of the 5-year Corporate Integrity Agreement to which Purdue was bound as a result of its 2007 guilty plea, when Rosetta Marketing Services LLC entered into a Master Services Agreement to work on OxyContin and other Purdue opioids.³⁹⁸ The Purdue relationship lasted as late as 2019.

555. As one industry trade journal described the relationship, "Rosetta's unique role lies in developing personalized marketing program built on consumer insights, as in the agency's integrated campaigns for... Purdue's pain drug OxyContin and pain patch Butrans" ³⁹⁹

1. OxyContin

556. From the outset of the Purdue relationship in 2010, Publicis entities, beginning with Rosetta, were Covered Persons⁴⁰⁰ pursuant to the Corporate Integrity Agreement Purdue was then bound by, and which remained in effect until May 2013.

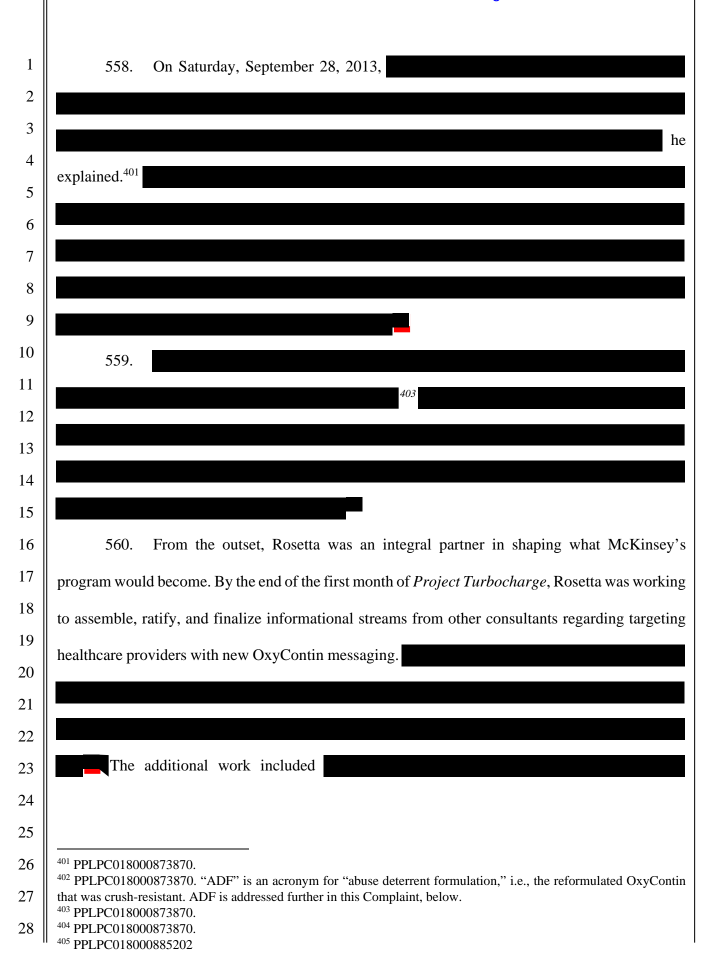
a. Project Turbocharge a/k/a Evolve to Excellence a/k/a E2E

557. Within a few *months* of its expiration, and with the CIA now out of the way, McKinsey proposed, and Purdue adopted, *Project Turbocharge*, a sweeping effort to revitalize OxyContin sales by overhauling and empowering Purdue's sales force to deliver in a precision-targeted way new messaging regarding OxyContin. Purdue adopted McKinsey's recommendations, coordinated with McKinsey to implement the recommendations, and immediately involved Rosetta in doing so. As a sign of the transformative nature of the undertaking, the project was unveiled as the theme of Purdue's 2014 national sales campaign.

Rosetta for \$575 Million," *New York Times*, May 17, 2011, *available at*: https://www.nytimes.com/2011/05/18/technology/18iht-publicis18.html
399 Marc Iskowitz, "100 Agencies: Rosetta – Acquisition by Publicis, integration activities lead to flat year for agency,"

Medical Marketing and Media, July 1, 2012, available at: https://www.mmm-online.com/home/channel/features/100-agencies-rosetta/

⁴⁰⁰ The relevant language in the Corporate Integrity Agreement provides: "'Covered Persons' includes . . . all contractors, subcontractors, agents, and other persons who perform sales, marketing, promotional, pricing, government contract, or regulatory functions . . . on behalf of Purdue."



561. Internally, Rosetta envisioned itself as perhaps playing an *even more* central role in McKinsey's *Project Turbocharge* at Purdue than it actually did. Rosetta described itself in an internal presentation as the "strategic backbone" of Purdue's marketing efforts, including *Evolve to Excellence*, which Rosetta identified as one of the definitive characteristics for "success" at Purdue in 2014. ⁴⁰⁷

562. Whether backbone or limb, Rosetta was present at the creation.⁴⁰⁸ The following sections describe certain components of Rosetta's work on *Evolve to Excellence* (or *E2E*, f/k/a *Project Turbocharge*). The conduct described is not exhaustive of Rosetta's (or Publicis') work in conjunction with McKinsey and for Purdue relating to *E2E* and adjacent projects. It is merely illustrative.

b. Titration and Length of Therapy

563. If you sell drugs to people, there are two ways to sell more drugs. One is to start selling the drug to someone who wasn't using it before (or, to use the lingo, a "new patient start"). The other way is to sell more of it to the people already using it. Titration and Length of Therapy ("LoT") efforts are directed at the latter opportunity. Stronger pills made more money; the higher the dosage strength for any individual OxyContin prescription, the greater the profitability for Purdue. Publicis went to great lengths to quantify the money to be made from increasing the length

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⁴⁰⁷ McKinsey's name, "Project Turbocharge" was seen as perhaps too crass or off-tone by Purdue. At a minimum, it was not "permanently appropriate." CEO John Stewart wrote to McKinsey partners Rob Rosiello and Arnab Ghatak on August 15, 2013: "Paolo Costa was especially engaged in the discussion and he (among others) will be a champion for our moving forward with a comprehensive 'turbocharge' process – *though we do need to find a better and more permanently appropriate name.*" (emphasis added). They settled on the decidedly more anodyne "*Evolve to Excellence*," (or "*E2E*").

⁴⁰⁸ See Dean Acheson, Present at the Creation: My Years in the State Department, W.W. Norton, (1969).

of therapy. According to their calculations, an increase of 1% in average LoT for yearly unique patients represents \$20 million in additional revenue to Purdue:

From: Ben Meck <ben.meck@rosetta.com> Date: Tuesday, August 19, 2014 at 11:09 PM To: John Dwyer < john.dwyer@rosetta.com> Subject: RE: ERO/OxyContin slide deck Okay, I can adjust based off this. · Which do you like better? Since you might be presenting o Every increase of 1% for average LoT for yearly unique patients (from 125.3 days on therapy to 126.6 days on therapy) is \$20M potential o Every increase of 1 day for average LoT for yearly unique patients (from 125.3 days on therapy to 126.3 days on therapy) is \$16M potential · Money point that I can include on the slide o Increasing average LoT by 2.5% for yearly unique patients (from 125.3 to 128.4 days on therapy) is \$50 million Ben Meck Manager | Analytics & Optimization Office +1 347.332.7655 Mobile +1 917.754.0041 99 Hudson Street, 11th Floor, New York, NY 10013, USA Rosetta.com Rosetta. Unlock and Activate™ Human Behavior.

- 564. Of course, higher dosage strength and increased lengths of therapy also contribute to opioid dependency, addiction and abuse. But Publicis was there to focus on selling higher strength dosages of OxyContin, and ROI was what was most important.
- 565. From 2012 through 2014, Publicis worked on numerous projects to design or refresh marketing campaigns to drive higher dose prescribing, for longer periods of time. In August of 2012, Publicis explained, "A strategic driver for [OxyContin] in 2013 is to drive appropriate titration and length of therapy with continuing patients. In an effort to add more emphasis to the importance of titrating to adequate analgesia... the brand team would like to create a 'campaign' to raise awareness."
- 566. Remarkably, Publicis created two *separate* marketing campaigns: one internal, for the Purdue sales representatives to understand how important titration and length of therapy is to *Purdue* (i.e., how the messaging effects profits), and a separate one designed to deliver the broader message to prescribers.
- 567. In September 2013, Publicis was brought in to "refresh" the already existing *Individualize the Dose* titration campaign. The campaign had its origins around four years prior

when, on October 26, 2009, McKinsey advised the Sacklers and the Purdue board that Purdue should train its sales representatives to "emphasiz[e] the broad range of doses," which would have the intended effect of increasing the sales of the highest (and most profitable) doses of OxyContin. McKinsey and Purdue subsequently implemented the campaign, and Publicis was brought in to do the "refresh" of the campaign in light of "an emerging market dynamic:" the decline in "mean patient dose" of OxyContin.

568. In other words, patients were buying less drugs. Publicis was brought in to reverse the decline; to "shift" the "trend":⁴⁰⁹



569. With respect to length of therapy, Publicis offered up a blunt instrument: coupons. In April of 2013, Publicis deployed an email marketing campaign, which sent out "Savings Cards" for OxyContin that could be downloaded. These coupons were known to be an effective method of encouraging patients to continue taking OxyContin for longer than they typically would otherwise.

- 570. The work continued into 2014 as E2E was being implemented, by which time Publicis was asking probing questions like, "Does a titration step lead to longer LoT?" "Does 1 titration step correlate with a higher likelihood of a 2^{nd} titration step? And a 2^{nd} to a 3^{rd} , and so on?"
- 571. In 2017, Publicis employees discussing titration messaging knew the score. As one put it, "We know discontinuation is usually an irrelevant subject matter (the persistent mindset is, once on an ERO [Extended-Release Opioid], the only way is up."
- 572. Higher doses of opioids taken for longer periods carry greater risk. Patients receiving high doses of opioids (e.g., doses greater than 100 mg morphine equivalent dose ("MED")

⁴⁰⁹ See 2013-09-03 Rosetta Creative Brief, cited in the Massachusetts Attorney General Complaint, Paragraph 57.

per day) as part of long-term opioid therapy are three to nine times more likely to suffer overdose from opioid-related causes than those on low doses. As compared to available alternative pain remedies, scholars have suggested that tolerance to the respiratory depressive effects of opioids develops at a slower rate than tolerance to opioids' analgesic effects. The Centers for Disease Control and Prevention also recognize that higher doses of opioids tend to increase overdose risks relative to any potential patient benefit.⁴¹⁰

- 573. Claims that opioids could be taken in ever-increasing strengths to obtain pain relief, without disclosing that higher doses increased the risk of addiction and overdose, are not only deceptive and misleading, they are deadly. These claims were particularly important to promotional efforts, however, because patients on opioids for more than a brief period develop tolerance, requiring increasingly high doses to achieve pain relief. Marketers needed to generate a comfort level among doctors to ensure the doctors maintained patients on opioids even as the high doses became necessary.
- 574. Publicis was ever vigilant in its protection of the titration messaging. In a June 2014 internal "Brand Overview" presentation regarding OxyContin, a Publicis employee struck out language in a draft version referring to utilizing the "lowest possible dose" because that language was contrary to the titration and length of therapy goals for the brand.
- 575. The vigilance paid dividends; the titration messaging worked. Nationwide, based on an analysis by the *Los Angeles Times*, more than 52% of patients taking OxyContin longer than three months are on doses greater than sixty milligrams per day, which converts to the ninety MED that the CDC guideline urges prescribers to "avoid" or "carefully justify."⁴¹¹

c. Targeting Patients; Targeting Prescribers

⁴¹⁰ Dowell D, Haegerich TM, Chou R. CDC Guideline for Prescribing Opioids for Chronic Pain – United States, 2016. MMWR Recomm Rep 2016;65(No. RR-1):1–49. DOI: http://dx.doi.org/10.15585/mmwr.rr6501e1

576. In order to aid prescribers and to deliver the titration message, Rosetta sought to differentiate and segment patients into different "types," so that physicians might familiarize themselves of instances in which titration is appropriate. In June 2014, in furtherance of the *E2E* initiative, Rosetta was hired to refresh two and create a third "patient profile," which could be left behind in brochure form in healthcare providers' offices.

577. These profiles were meant to personalize OxyContin patients in the eyes of the prescriber and described the instances in which an OxyContin patient's dosage should be changed. "Maggie" is 43 years old with "significant" lower back pain. After falling at home, she begins a 40mg OxyContin twice daily regimen, and is then titrated up to a 60mg tablet twice daily before she "reports that her pain is properly managed." "Carol" is 51 years old and has osteoarthritis in her left hip. She starts at a 15mg dose and is titrated up to 20mg dose, "with better relief of pain symptoms." "James" is 40 and his osteoarthritis is in his knee. He starts at a 10mg OxyContin dose, then is doubled titrated to 15mg. After reporting that "his pain is still not well managed," James is titrated up again to a 20mg dosage, at which point he experiences "good response to his arthritic pain."

578. Notably, in none of these patient "vignettes," as Rosetta called them, was a patient ever titrated *down*. With EROs like OxyContin, the goal of titration was always ever skyward. These patient vignettes were to be included alongside information about obtaining "savings cards" – coupons – in order to get a discount on your OxyContin prescription. The savings cards were offered because Publicis knew from experience that their use correlated with longer average length of therapy for the patients that used the coupons.

579. Another patient group that Rosetta targeted were individuals already prescribed short-acting opioid medications. The goal was to transition these patients to an extended-release

formulation for long-term management of their pain. A longer length of therapy on an ERO like OxyContin is more profitable than sales of short acting opioids.

- 580. As Rosetta segmented and targeted patient types, so too did it segment and target certain prescribers. It did so in order to deliver tailored messaging to targeted individuals in order to maximize the goal, to
- 581. As mentioned above, from the outset of *E2E*, Rosetta was working on identifying target lists of OxyContin prescribers,

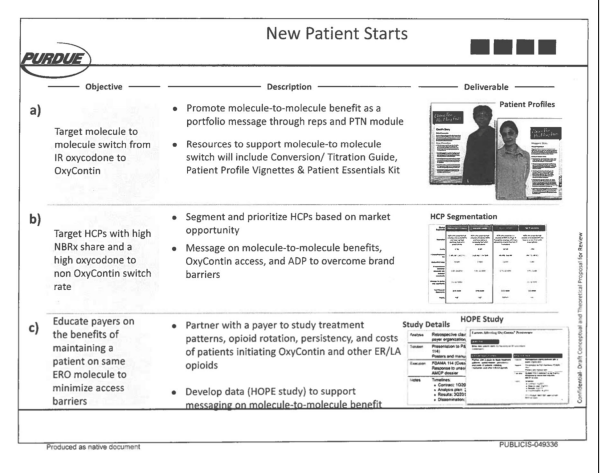
Rosetta's targeting efforts were organized by segmenting OxyContin prescribers into deciles by volume of prescriptions written.

- 582. Once these target lists were compiled, the purpose was to concentrate marketing efforts on the highest decile prescribers, i.e., those doctors who were already prescribing OxyContin and who, upon information and belief, had previously been subjected to Purdue's tainted marketing efforts *prior* to the 2007 guilty plea.
- 583. At the same time, Publicis also focused marketing efforts on *new* prescribers who were prescribing more OxyContin than their peers. This separate market segment of "new to brand prescribers" (or "NBRx") could be plumbed for additional growth. Thus, Publicis advised sending sales representatives to both "high decile prescribers" and "High-decile NBRx HCPS" up to three times a month. These efforts were part of an overall initiative to generate "new patient starts" (which is one way to increase sales, the other being Titration and Length of therapy, described above), as set forth in an "OxyContin 2015 Tactical Planning" presentation prepared by Rosetta:

- 153 -

⁴¹² PPLPC018000873870.

⁴¹³ PPLPC018000885202.



- 584. Rosetta identified these physician targeting efforts to sell more OxyContin, which could result in more than \$1 billion of additional revenue for Purdue.
- 585. The targeting was not limited to segmenting physicians into deciles by volume of OxyContin prescriptions written. It also identified different physician profiles. Two patient profiles in particular were identified for targeting. The "motivated believer" was "convinced that opioid medication is essential to treatment of the chronic pain patient" and recognized OxyContin as "the leading FDA-recognized Opioid with Abused Deterrent Properties." A "brand loyalist" was a prescriber that would remain an OxyContin prescriber despite growing concern within the medical profession regarding its abuse liability.
- 586. One difficult physician segment type is the "no-see." Some physicians simply do not want to see sales representatives and will not meet with them in person. Obviously, this creates an impediment to access that pharmaceutical manufacturers seek to surmount. If you cannot

communicate to a physician, it is difficult to influence that individual's prescribing patterns. Rosetta offered solutions.

- 587. Of course, there are other channels that can be used, such as email marketing campaigns, and this Rosetta did. But there are other ways to gain access. For instance, Publicis suggested that a list of all "no-see" prescribers who worked at "integrated delivery networks" be compiled: "We should target anyone on the IDN Lists… regardless of decile." Rosetta and Purdue then set up a call center where these prescribers who did not wish to meet with sales representatives would receive phone calls instead.
- 588. Another response to barriers to physician access is to go around them, to talk to someone else at the doctor's office instead. Rosetta designed strategies to target physician assistants and nurse practitioners instead of the physicians themselves. Like the physician profiles it created, Rosetta also segmented the physician assistants and nurses into "attitudinal and behavioral profiles," the better with which to target OxyContin messaging, such as "dose titration opportunities." In 2015, Rosetta proclaimed that these efforts to target nurses and physician assistants could yield an additional 14,500 OxyContin prescriptions in the first year of implementation.
- 589. Two years later, Publicis' efforts at targeting nurses and physician assistants continued, with a Publicis employee noting the particular importance of this market segment for OxyContin. In a September 27, 2016, email, she noted, "NP/Pas are a growing provider segment and help offset the decline from PCPs."
- 590. Publicis also saw their work on targeting nurses and physician assistants as a cross-selling opportunity. In 2016, Rosetta suggested Purdue hire Rosetta's affiliate, Publicis Touchpoint Solutions, to deploy clinical nurse educators "trained on the Purdue sales force platform" to help

of the ov

these nurses and PA's "understand and implement" OxyContin treatment plans and "overcome behavioral obstacles that may interfere with patient adherence."

591. As a part of the overall implementation of *E2E*, Rosetta was involved not only in delivering the marketing messages that Purdue would then deliver to healthcare providers. Rosetta remained deeply involved in the actual rollout of these marketing messages. In July of 2015, three Rosetta employees on the Purdue account joined sales representatives in the field for ride-alongs to meet with targeted prescribers. These Rosetta employees observed in person the interactions between Purdue sales representative and target prescribers during which Rosetta's messaging was delivered. Rosetta was able to use the knowledge they gained from observing prescribers' reactions to their messaging, which further refined their approach and maximized the effectiveness of the marketing outreach.

592. These field trips served the overarching goal, as Purdue's Ron Cadet explained, to

d. Non-indicated Uses

593. Publicis introduced a search engine marketing "Conditions Campaign" for OxyContin as early as 2013 and continued these efforts throughout the implementation of *E2E*. The campaign was designed to drive prescribers conducting web searches about certain medical conditions – lower back pain, for instance – to Purdue's website. "The Campaigns pertaining to Conditions were the most successful out of last year's [search engine marketing] strategy and

⁴¹⁴ Consistent with the "Power of One" approach described supra., this was not the first time Publicis cross-sold other Publicis divisions into the Purdue account. In 2015, Razorfish advised Purdue to hire Verilogue without going through a competitive bidding process. This cross-selling initiative – a fine example of the "Power of One" in action – resulted in Verilogue's analysis of patient-provider conversations being used to devise to tactics to increase sales volumes of Purdue's opioids. Verilogue's was used in furtherance of targeting and messaging goals of *E2E*, and its work was part of the overall efforts to

⁴¹⁵ PPLPC018000873870.

resulted in 25,548 Clicks, driving the most visits to PurdueHCP site," Publicis explained in a June 2014 presentation.

594. The Conditions that Publicis targeted included low back pain, cancer pain, and osteoarthritis pain. OxyContin is only approved, or indicated, to treat certain conditions, and is not approved to be marketed for a non-indicated used.

595. Publicis proposed to expand the Conditions campaign to target multiple sclerosis, trigeminal neuralgia, and spinal cord injury, but the proposal "raised some red flags." John Dwyer explained in an internal Publicis email, "there are search words that are disease states OxyContin is not indicated for. And by having it in a document associated with OxyContin even for internal review can put [Purdue] in a liability risk." Accordingly, Publicis altered the text of the advertisements that would appear when a given disease state was searched for in order to delete reference to that disease state. (The ads still appeared when prescribers searched for the non-indicated Conditions, the text of the ad merely referred to "that" condition, instead of a specific reference to multiple sclerosis, low back pain, or the like.)

2. Purdue's other brands: Butrans, Hysinglia, et. al.

596. While OxyContin accounted for the lion's share of Purdue's revenue, the monoline manufacturer sold other flavors of opioids. By the end of 2016, Publicis had won the agency of record position for the rest of the Purdue's brands as well.

597. In a 2014 presentation,

|| 416 PPLPC012000477940

Rosetta has a long standing history with Purdue across multiple brands

2010 - Started as OxyContin Agency of Record for HCPs (as Wishbone)

- AOR for Partners Against Pain (PAP)
- RM program and Portal pilots for OxyContin
- Marketing Education projects
- Purdue corporate brand campaign development, corporate website, and other corporate materials





2011 - RM Management for launch of Butrans

PurduePharma.com work, digital media planning for OxyContin



2012 - RM Management for launch of Intermezzo

- Classwide Opioid REMS website working in partnership with McKesson (ER-LA-opioids,com)
- Purdue products' REMS website (PurduesREMS.com)



Intermezzo

(ZOLPIDEM TARTRATE) sublingual tablet @

1.75 mg | 3.5 mg

2013 – Won the AOR Assignment for Targinia ER

Multichannel ADF Campaign evolved from prelaunch planning



2014 - RM Management and Digital Media for HYD

Publicis worked on Purdue's Butrans, Intermezzo, Targiniq, and Hysinglia brands in addition to OxyContin.

598. The same year, John Dwyer met with Purdue's Peter Justaston to discuss "Agency consolidation," and what Publicis "would need to do" to win the single Agency spot. Dwyer assured Purdue that Publicis was "up to the challenge" of being the sole Agency for all of Purdue's products. It was a goal Publicis pursued doggedly.

599. "For the last 4 years, every year as we do the forecast for the new year, I've put 'Butrans AOR business' in there under whitespace, which is where we put areas where we think we can grow," John Dwyer said to his team at Publicis in April 2016. That year, Publicis achieved Dwyer's goal, with Purdue deciding to shift some work it previously assigned to competitor agencies to Publicis' Razorfish.

600. Dwyer sent a congratulatory email on April 4, 2016, to the Publicis team about Publicis' winning additional business from Purdue, and explained that their success on the OxyContin account is what drove the decision for Purdue to award Publicis' Razorfish additional work:

... when the client told us that he was going to move the rest of the Butrans business over to Razorfish Health, the reason he said he was doing it was because he wanted up to replicate exactly everything we did last year [2015] on OxyContin. The creative campaign refresh, the vis[ual] aid that far outshone the Butrans and Hysinglia ones which were done by another agency, the strategic planning we brought them for the new competitors who are coming, and the overall revitalization of the brand... He said, "We want more like that. We want you to do the exact same thing this year, but now with Butrans.

601. The following month, Razorfish was awarded the Hysinglia account as well. Publicis' Karl Tiedemann noted in an email the "amazing relationship" developing between Purdue and Publicis, and quoted a Purdue employee as stating that the Hysinglia account was "the final piece and we now own Purdue." (emphasis added).

3. Selling red herring: ADF

- 602. One way to keep selling opioids once widespread abuse and dependence has gripped the patient population that uses them is to introduce a new version of the drug that can be sold as "abuse deterrent." The opioid manufacturers did just that, with Purdue leading the way. Publicis was there at every step of the way to define the message and deliver it to crucial audiences: prescribers, patients, and regulators.
- 603. Purdue's "[abuse deterrent formulation] does not prevent abuse via swallowing," Publicis stated in a June 2014 confidential internal memorandum.
- 604. "As long as a drug is addictive, can be abused by swallowing higher doses, and ways to defeat the [abuse deterrent formulation] are online," no drug would ever meet the FDA's standard for "reduced abuse, misuse, and/or diversion in the community,"

explained former Publicis partner Ted Whitby in a June 15, 2017, email to his former colleagues at Publicis.

605. In other words, the new formulations didn't prevent abuse. They were red herrings, a distraction that an opioid manufacturer could point to in order to mollify regulators and other interested parties, while knowingly not addressing the underlying abuse liabilities of their products. Publicis knew this good and well, but never mind that. ADF was a message to deliver.

606. And Publicis was eager to spread the word. In July 2014 (*one month* after the internal Publicis memorandum discussing the ineffectiveness of the abuse-deterrent formulation), Publicis prepared the following language as a "value proposition" that sales representatives could recite to doctors: "OxyContin is the only opioid that addresses the pain management needs of both patients (efficacy and dosing flexibility) and society (proven abuse deterrent properties)."

607. This work was part of an ongoing project by Rosetta to get the ADF message out to the world for Purdue. On March 31, 2014, as part of the overall *E2E* initiative,

was misleading, but craft it they did, nonetheless. In May 2015, Publicis was working on creating

The work proved grim. Publicis knew that the ADF messaging they were crafting

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⁴¹⁷ PPLP0033598595

⁴¹⁸ PPLP0033598595

⁴¹⁹ PPLP004128540

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609. In response to the growing crisis, in March 2016 the Centers for Disease Control and Prevention issued a Guideline for Prescribing Opioids for Chronic Pain (the "CDC Guidelines") in order to reduce the recognized overprescribing of opioids and the concomitant rise in opioid use disorder.

- 610. Within a month of their issuance, Publicis provided an assessment of the threats and opportunities that each of the twelve Guidelines posed. The "threat" was that the Guidelines would result in reduced OxyContin prescribing. The "opportunity" was that the Guidelines could be used to position Purdue's products to sell *more* of them by *using* the Guidelines.
- 611. In general, the Guidelines were seen as a threat as they recommended actions that ran directly counter to Purdue's marketing efforts. For instance, the Guidelines recommended that prescribers use the "lowest effective dosage" when prescribing opioids, which is anathema to the titration and length of therapy messaging that Publics and Purdue had been articulating and delivering for years.
- 612. The Guidelines also called for a curtailment of prescriptions exceeding 90 morphine milligram equivalents per day. These high dose prescriptions accounted for nearly half of total OxyContin prescriptions and were each more profitable than their lower dosage equivalents. The Guidelines therefore risked to substantially erode OxyContin revenue. Purdue estimated the cost to Purdue resulting from implementation of the Guidelines could be in excess of twenty million dollars annually in lost sales in single state.

iii. **Unbranded Marketing**

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- It might be obvious, but "unbranded" opioid marketing does not promote any 613. particular brand of opioid. Instead, unbranded marketing promotes opioids as a *class* of drugs worthy of ever increased prescription. Here, Publicis excelled in its work for Purdue.
- 614. If one thinks of the entire opioid market as a pie, each individual brand (as well as generic formulations) would comprise individual slices. Crucially, even if an individual product's market share does not change, that portion can grow in value if the *overall* market grows. The larger the pie, the larger the slice.
- The long-term goal was to "make the whole pie bigger, not only for us, but for our competitors as well," a Purdue executive explained in a 2000 speech. 420

1. **Practical Tools to Advocate for yourself.**

616. On January 22, 2012, Jennifer Grey, the actor known for co-starring in "Dirty Dancing," appeared as a "patient advocate" on a local television talk show in San Francisco to discuss "Partners Against Pain," a campaign with which she was affiliated. Noting the website's address for the audience, she explained, "it's called PartnersAgainstPain.com, and it's a really interesting educational program... It's a campaign that was meant to help people become advocates for themselves who are in chronic pain and because chronic pain is such an insidious syndrome."⁴²¹

available

⁴²⁰ Sari Horwitz, Scott Higham, Dalton Bennet and Meryl Kornfield, "Inside the opioid industry's marketing machine," Washington Post. December https://www.washingtonpost.com/graphics/2019/investigations/opioid-marketing/

⁴²¹ Sari Horwitz, Scott Higham, Dalton Bennet and Meryl Kornfield, "Inside the opioid industry's marketing machine," Washington Post. December 2019, available https://www.washingtonpost.com/graphics/2019/investigations/opioid-marketing/



617. In 2019, Grey provided the *Washington Post* with the following statement regarding her prior affiliation with PAP⁴²²:

When this unbranded [Purdue] Pharma campaign was brought to me in 2011, I was excited by the opportunity to help people who, like myself, suffered from chronic pain by giving them practical tools to advocate for themselves and manage their pain in a safe, responsible way. I never suspected I was being used to potentially advance a darker agenda. (emphasis added).

618. Rosetta and Purdue designed the PAP campaign together.

They were the ones who "brought" the "unbranded [Purdue] Pharma campaign" to Grey.

619. From the outset,

⁴²² Sari Horwitz, Scott Higham, Dalton Bennet and Meryl Kornfield, "Inside the opioid industry's marketing machine," *Washington Post*, December 6, 2019, *available at*: https://www.washingtonpost.com/graphics/2019/investigations/opioid-marketing/

⁴²³ PPLP003518651

⁴²⁴ PPLP003518651

(emphasis added).

623. In other words, Grey was hired to sell more drugs. Her work was just another line item in the marketing budget, and Rosetta and Purdue expected a return on the money spent for it.

624. Not content with Grey alone, Rosetta and Purdue enlisted Country Music Icon Naomi Judd as well. "Judd wants people to know the journey to appropriate pain management can begin with a visit to the recently updated Partners Against Pain website (www.PartnersAgainstPain.com)," proclaimed the press release. 432

625. Like Grey, Judd appeared in television spots as a "patient advocate": 434

434 See https://www.youtube.com/watch?v=cUav8M9mep4

⁴³¹ PPLPC017000328575, at 328587.

⁴³² See https://ftper.newsusa.com/Pdfs/NaomiJudd.pdf

⁴³³ PPLP003518651 at 3518686 (underlined emphasis in original; italicized emphasis added –



- 626. "From day to day, pain can limit your ability to work, your hobbies, even the simple joys of huggin' somebody you love," she explained. 435
- 627. Five years later, in 2016, Publicis proposed to revamp the *Partners Against Pain* website in light of the fact that the website had been "named in lawsuits." Better to "start from scratch," Publicis proposed. Publicis instead pitched to Purdue its "Patient Support Program." The ultimate goal of the new campaign would be the same as the original *Partners Against Pain* campaign, "extending patients' length of therapy."

2. Join the Team!

628. Razorfish Health was the agency of record for Purdue in connection with the launch of the now-defunct website TeamAgainstOpioidAbuse.com. 436 Purdue claimed the website was

⁴³⁵ See https://www.youtube.com/watch?v=cUav8M9mep4

⁴³⁶ Kevin McCaffrey, "Purdue debuts opioid-abuse resource," Medical Marketing and Media, August 17, 2015m, *available at*: https://www.mmm-online.com/home/channel/campaigns/purdue-debuts-opioid-abuse-resource/

"designed to help healthcare professionals and laypeople alike learn about different abuse-deterrent technologies and how they can help in the reduction of misuse and abuse of opioids."

629. The unbranded website contained misleading information regarding the effectiveness of abuse-deterrent properties of certain opioid formulations, including Purdue's reformulated OxyContin and Hysinglia. At the time, only three abuse-deterrent formulations were available on the market (the third was Embeda, manufactured by Pfizer). The concept is that we're all part of a team," explained Dr. David Haddox, Purdue's Vice President of Health Policy. 439

3. The Meaning of the Message

- 630. There was a recursive nature to the marketing of opioids as companies evolved from *growing* the market for opioids to *responding* to the abuse and dependency issues that were the natural outgrowth of that initial boom in opioid sales. "Hello, my name is Mark Timney... We at Purdue are committed to doing everything we can to reverse this public health problem," read the script written by Rosetta for newly-installed Purdue CEO Mark Timney to read on a 60-second television ad prepared by Rosetta in 2014.
- 631. Initially, opioids were marketed primarily based on their efficacy (while disregarding their abuse liability). The PAP campaign was one facet of that effort. Once abuse became a concern, opioid marketers then sought to contextualize opioid addiction within a broader context of the individuals' right to manage their own pain as a part of their own interactions with the nation's healthcare system. The words Timney said on TV and the website TeamAgainstOpioidAbuse.com exemplify this later approach.

^{437 &}quot;Purdue Pharma L.P. Launches TeamAgainstOpioidAbuse.com," Purdue, Aug. 17, 2 http://www.purduepharma.com/news-media/2015/08/purdue-pharma-l-p-launches-teamagainstopioidabuse-com/.

⁴³⁸ Kevin McCaffrey, "Purdue debuts opioid-abuse resource," Medical Marketing and Media, August 17, 2015m, *available at*: https://www.mmm-online.com/home/channel/campaigns/purdue-debuts-opioid-abuse-resource/
⁴³⁹ *Id*.

- onstructed around the theme of abuse prevention a decision that works strategically to shield the company [Purdue] from the negative press and attention being directed at its products." ⁴⁴⁰ But while the website did not specifically name brands of abuse-deterrent opioids, the website nonetheless operated to promote further use of Purdue's drugs. Of the three abuse-deterrent opioid products then on the market, Purdue manufactured two of them. As such, "the site also functioned as a marketing platform for those same drugs, which it framed as a solution to the problem of opioid addiction and abuse."
- 633. It is no accident that Ms. Grey spoke about providing "practical tools" to *consumers* in order to allow them to "advocate" for themselves. These tactics, subtle and insidious as they are, are meant to increase overall opioid prescribing. "Pharmaceutical companies often mobilize education campaigns around a particular diagnosis in order to indirectly market the cure and as a mean of evading legal constraints that restrict the ways that companies advertise their drugs to consumers."
- 634. Ms. Grey expressed concern about being used for a "darker agenda." It was simply to sell more drugs.
- 635. For example, the *Partners Against Pain* website that Ms. Grey promoted "groomed" potential opioid customers by offering assessment tools to website visitors including "checklists and lists of questions used to guide patients through the process of obtaining medication from their

 $^{^{440}}$ Melina Sherman, "How to Train Your Opioid Consumer," $\it Communication, Culture \& Critique Vol. 10 (2017), Pgs. 600-601.$

⁴⁴¹ *Id*.

⁴⁴² Melina Sherman, "How to Train Your Opioid Consumer," *Communication, Culture & Critique* Vol. 10 (2017), Pg. 599 (citing Angell, M. (2004). *The truth about drug companies: How they deceive us and what to do about it.* New York, NY: Random House).

doctors," as a means of educating the viewer. 443 The website also included a "Find a Doctor" tool, whereby viewers could search for pain management practitioners in their local area.

- 636. As the Rosetta ad that Purdue CEO Mark Timney read on TV emphasized, "Purdue is greatly expanding our ongoing efforts to help educate the public about prescription opioid abuse. We will be doubling our investment in 2015 to continue building awareness about the problem and better educating physicians, pharmacists, school health officials, insurers, and lawmakers about the risks of opioid abuse." TeamAgainstOpioidAbuse.com was one such educational effort.⁴⁴⁴
- 637. But "tools such as these go further than education: They also function as technologies of power that manage communication and information in a way that is normative and directional and, in so doing, guide visitors' behavior toward actions that benefit the company."
- 638. Dr. Marcia Angell, former Editor in Chief of the *New England Journal of Medicine*, summed things up plainly:⁴⁴⁶

No one should rely on a business for impartial evaluation of a product it sells. Yet the pharmaceutical industry contends it educates the medical profession and the public about its drugs and the conditions they treat, and many doctors and medical institutions – all recipients of the industry's largesse – pretend to believe it. So does the government. But "education" comes out of the drug companies' marketing budgets. That should tell you what is really going on.

639. Publicis employee Karl Tiedemann summed it up tidily, stating that the work Publicis was doing for Purdue was "not about education, but persuasion."

⁴⁴³ Melina Sherman, "How to Train Your Opioid Consumer," *Communication, Culture & Critique* Vol. 10 (2017), Pg. 599.

⁴⁴⁴ Even today, these educational efforts are cited as justification for Purdue's actions. See Eric Russell, "Portrayed as villain in TV series about opioid crisis, ex-U.S. attorney for Maine says he didn't sell out," Portland Press Herald, February 27, 2022, available at: https://www.pressherald.com/2022/02/27/portrayed-as-villain-in-tv-series-about-opioid-crisis-ex-u-s-attorney-for-maine-says-he-didnt-sell-out/ ("McCloskey said the scene depicts an inaccurate timeline and he was especially disappointed with the insinuation that he was paid by Purdue to keep quiet. 'There's no truth to that whatsoever,' he said. 'Purdue did a number of things I asked them to do. They took a pill off the market because I asked them to do it. They worked on tamper-proof prescription pads. They spent millions producing educational brochures.'") (emphasis added).

⁴⁴⁵ Melina Sherman, "How to Train Your Opioid Consumer," *Communication, Culture & Critique* Vol. 10 (2017), Pg. 599.

⁴⁴⁶ Dr. Marcia Angell, *The Truth About the Drug Companies*, Random House, 2004, Pg. 135.

640. A 2015 Publicis presentation backed up Tiedemann's statement. It identified "patient education" and "increase[ing] awareness as the "strategies" to achieve the "objectives" of "grow[ing] volume" in light of, among other things, "increasing pressure to reduce abuse, diversion, and overdose:"

2015 OxyContin Strategic Imperatives Updated 7/10		
Opportunity/Issue	Objective	Strategy
No IROs have ADPs Likely national up-scheduling of hydrocodone Segmented messaging with a	Grow volume/ slow erosion	Increase conversions from oxycodone IROs by educating HCPs on how to identify appropriate patients
promotionally-sensitive brand Little approvable branded messaging to drive IRO-to-ERO conversion Continued declining mean patient dose Patient understanding of chronic pain management		Drive appropriate titration through ongoing reassessments and patient education
Increasing pressure to reduce abuse, diversion and overdose Limited prescriber understanding of current OxyContin ADP evidence Potential Tier 4 approval	Increase choice of OxyContin as 1 st branded ERO through meaningful differentiation	Increase awareness, understanding and relevance of OxyContin abuse deterrent properties to drive preference vs. all other opioids
Potential approval of new ER oxycodones with ADPs OxyContin has 7 tablet strengths Continued pharmacist role in opioid Rx vetting		Maintain payer coverage; maximize pull-through opportunities and patient affordability

641. Moreover, unbranded marketing schemes like the one in which Ms. Grey unwittingly participated have a broader purpose of creating an information ecosystem designed to benefit the interests of pharmaceutical companies. As Verilogue noted, by "shaping the dialogue," one can "shape the future."

642. Dr. Sherman explained,

This practice of dispersing branding materials in new contexts is emblematic of what Dumit (2012) has referred to as "strategic ubiquity," a tactic in which companies attempt to create a "universe" of syndicated and sponsored content through forming alliances with advocacy groups and developing partnerships with other influential third parties. For potential customers navigating this landscape, *every piece of*

⁴⁴⁷ See infra, fn. 493 and accompanying text.

information they read or hear about inevitably directs them toward specific actions that will serve the benefit of the company.⁴⁴⁸

- 643. The strategic intent of these campaigns, then, is "to construct an echo-chamber of pro-opioid information." 449
- 644. One example of this echo-chamber is Publicis' groundbreaking work with *regulators* on their own opioid marketing campaigns. In 2018, Razorfish Health partnered with the FDA's Center for Drug Evaluation and Research (CDER) to design the "Search and Rescue" campaign, which "makes innovative use of optimized search and direct email to reach family physicians, physician assistants and nurse practitioners, and in states with the highest opioid prescribing rates." Razorfish's campaign for the FDA was designed to further the same educational messages Purdue was espousing, to "equip prescribers to be proactive in identifying and helping patients at risk for prescription drug abuse." 451
- 645. In other words, Publicis was able to deliver Purdue messages through alternative channels, including not only unbranded websites, but through the mouthpiece of *Purdue's regulator*, the United States Food and Drug Administration, as well. *That* is an echo-chamber.

iv. Qui audet adipiscitur

646. "By nature, healthcare advertising provides a crutch to advertisers and marketers in the form of industry guidelines and regulations. We're quick to point out why we *can't* do

⁴⁴⁸ Melina Sherman, "How to Train Your Opioid Consumer," *Communication, Culture & Critique* Vol. 10 (2017), Pg. 603. (*citing* Dumit, J. *Drugs for life: How pharmaceutical companies define our health*. Durham, NC: Duke University Press) (emphasis added).

⁴⁴⁹ Melina Sherman, "Opiates for the masses: constructing a market for prescription (pain)killers," *Journal of Cultural Economy*, Vol. 10, Issue 6, 2017, *available at*: https://www.tandfonline.com/doi/full/10.1080/17530350.2017.1352010

⁴⁵⁰ See https://drugfree.org/newsroom/news-item/partnership-drug-free-kids-announces-relaunch-search-rescue-opioid-prescriber-education-campaign-website/

451 Id.

risk and drive real innovation in healthcare marketing.

something, rather than eager to question how we could make something work within those boundaries," pronounced Publicis Health Media's Media Director, Eric Delash. 452

647. Instead, Delash encouraged pharmaceutical marketers to "embrace risk:"453 Challenge the status quo. Disrupt the norm. Create your Gritty. Embrace strategic

648. Delash's dashing attitude is reminiscent of prior rakish campaignors. "Who dares wins," chose Sir David Stirling before him. 454 Naturally, Publicis has long embodied the principles espoused in its Media Director's blog post. Indeed, Publicis has embraced risk and dared greatly⁴⁵⁵ on behalf of its clients. In 2016, Publicis submitted proposed work for the aforementioned TeamAgainstOpioidAbuse.com website, as well as a separate campaign for Hysinglia that it had pitched to Purdue, for consideration in the category of "Most Daring Campaign" in Medical Marketing and Media's annual industry awards. 456

Coda Part II v.

649. Success is determined by what is being measured. In one sense, Purdue was an inordinately profitable account for Purdue. Publicis' relationship grew from its start in 2010 to

⁴⁵² Eric Delash, Healthcare Marketers Must Embrace Risk to Innovate, Publicis Health Media Blog, December 13, 2018, available at: https://www.publicishealthmedia.com/perspective/healthcare-innovations/

⁴⁵³ Eric Delash, Healthcare Marketers Must Embrace Risk to Innovate, Publicis Health Media Blog, December 13, 2018, available at: https://www.publicishealthmedia.com/perspective/healthcare-innovations/

Stirling: See David The Phantom Major, National Army Museum, available at: https://www.nam.ac.uk/explore/david-stirling (recounting the origin of the motto of the British Special Air Service – Qui audet adipiscitur - during World War II); see also: https://en.wikipedia.org/wiki/Who Dares Wins

⁴⁵⁵ See another rakish campaignor, Theodore Roosevelt, supra, fn 16. ("It is not the critic who counts; not the man who points out how the strong man stumbles or where the doer of deeds could have done them better. The credit belongs to the man who is actually in the arena, whose face is marred by dust and sweat and blood; who strives valiantly; who errs, who comes short again and again, because there is no effort without error and shortcoming; but who does actually strive to do the deeds; who knows the great enthusiasms, the great devotions; who spends himself in a worthy cause; who at the best knows in the end the triumph of high achievement, and who at the worst, if he fails, at least fails while daring greatly, so that his place shall never be with those cold and timid souls who neither know victory nor defeat.")(emphasis added).

⁴⁵⁶ Incredibly, that year, "MM&M's esteemed and independent panel of judges elected not to award a Gold for Most Daring Campaign... While some excellent and creative unsold work was entered, the judges felt that none of the submissions were precisely 'daring' enough, according to the judging criteria, to receive the top honor." Tanya Lewis, Most Daring Campaign of 2016, Medical Marketing and Media, October 6, 2016, available at: https://www.mmmonline.com/mmm-awards/most-daring-campaign-of-2016/ Not all who dare, win.

encompass practically all branded marketing business from Purdue six years later. By 2016, Publicis was the agency of record for OxyContin, Butrans, Hysinglia, and Targiniq. "The growth we continue to see in this business is phenomenal," congratulated a Publicis group president to the Purdue team. On March 22, 2016, John Dwyer emailed Karl Tiedemann to let him know that the expected annual revenue from Purdue for that year was approximately \$12 million. "Oh boy," replied Tiedemann. The Subject line of the email thread was "We're gonna need a bigger boat." In that sense, Publicis was very successful.

- 650. Viewed differently, Publicis' daring work was outrageously successful for Purdue and, more to the point, the Sacklers. After 2013, the goal of *E2E* was to
- and this Publicis did with great success. The Sackler Strategy that Publicis sold yielded large amounts of money available for distribution from Purdue to the Sackler family for over a decade. In that sense, Publicis was very successful.
- 651. In another sense, Purdue itself ended up bankrupt and, just like its parent, the Purdue Frederick Co., before it, a convicted felon. This might be something other than success.
- 652. On October 20, 2020, Purdue—Publicis's co-conspirator—agreed with the United States Department of Justice to plead guilty to improper marketing of OxyContin and other opioids again (the "2020 Settlement Agreement"). This time the plea agreement concerned conduct from 2010 to 2018. The agreement includes \$8.3 billion in penalties from Purdue and \$225 million from the Sackler family.
- 653. Purdue pled guilty to a dual-object conspiracy to defraud the United States and to violate the Food, Drug, and Cosmetic Act, 21 U.S.C. §§ 331, 353, violating anti-kickback laws, and "using aggressive marketing tactics to convince doctors to unnecessarily prescribe opioids —

⁴⁵⁷ PPLPC018000873870.

frivolous prescriptions that experts say helped fuel a drug addiction crisis that has ravaged America for decades."

654. The new plea agreement did not identify Purdue's co-conspirators, but Purdue's new guilty plea concerned Covered Conduct (as defined in the plea agreement) that directly implicates Publicis in the conspiracy, including the same conduct described in this Complaint.

m. Publicis and Other Opioid Manufacturers

655. Publicis' role in the propagation of the opioid epidemic extends far beyond its work with felons. During the pendency of its long-term relationship with Purdue, and their joint efforts to grow the *overall* opioid market, Publicis also partnered with Endo, Teva, Janssen, and others to market those clients' opioid brands. While it sought to grow the overall pie, Publicis also endeavored to grow each slice. It attacked the problem from both angles. And in doing so, it deployed many of the same tactics, at the same time, for numerous competing opioid brands. If some of the following paragraphs appear redundant, there is a reason. Lots of manufacturers' opioid marketing strategies looked a lot alike. Publicis was a common denominator.

i. Endo

- 656. Publicis' Saatchi and Saatchi unit ("Saatchi") was the Agency of Record for Opana, Endo Pharmaceutical's branded oxymorphone product. As described above, Opana is the same molecule as Endo's previous product Numorphan depicted in the film *Drugstore Cowboy*.
- 657. With the launch of Opana, Endo decided it was time for history to repeat itself. After Opana's approval in 2006, Endo solidified its position as a pain specialist among manufacturers. By 2012, Endo's opioid sales accounted for approximately \$403 million of its \$3 billion in revenue, more than 10% of its total sales. From 2010 to 2013, total Opana ER revenue alone exceeded \$1.1 billion.

touted its work with co-Defendant Allscripts as a "main win." Id.

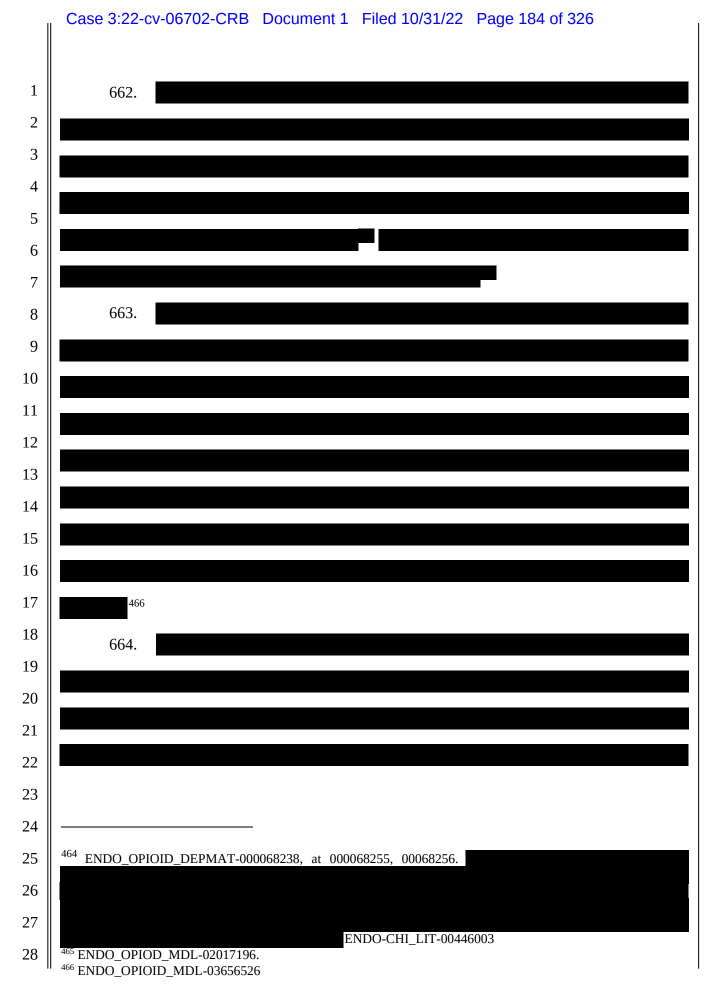
⁴⁶³ ENDO OPIOID DEPMAT-000068238, at 000068258.

⁴⁶¹ ENDO-OPIOID MDL-02090726

⁴⁶² ENDO FLAG-01075412

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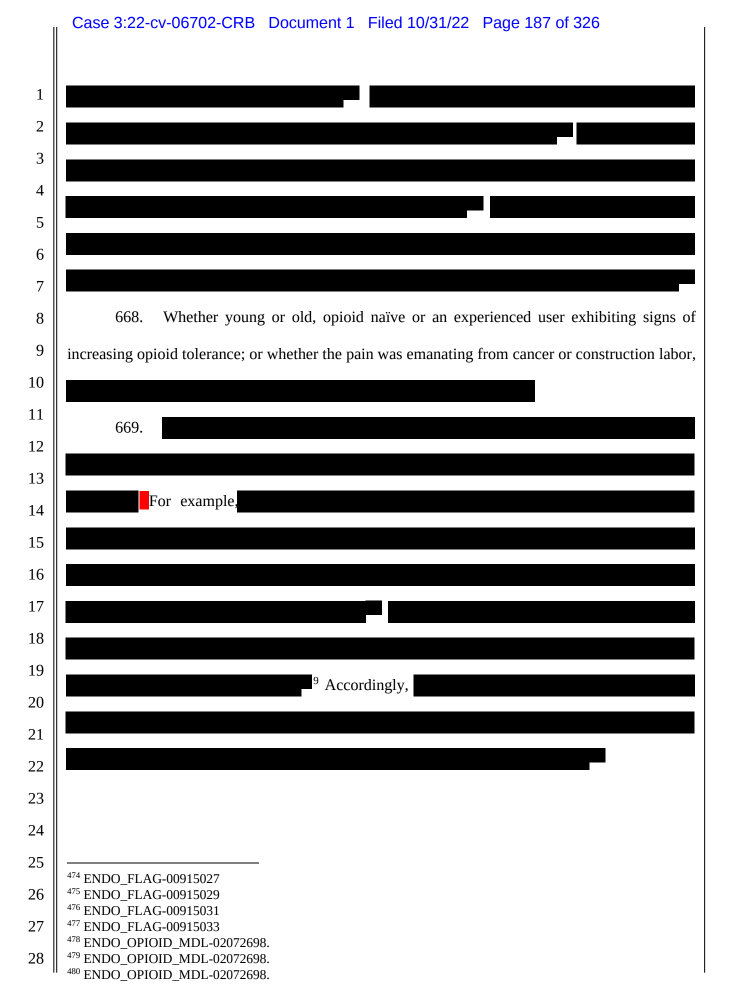


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⁴⁷² ENDO_FLAG-00911567

⁴⁷³ ENDO_FLAG-00915025



1 Also in 2007, 670. 2 3 Saatchi relied on the fact that the Joint Commission on Accreditation of Healthcare Organizations 4 (JCAHO) had recently adopted pain to as "the fifth vital sign," and now required hospitals to assess 5 and treat patients' pain as a matter of course. 482 JCAHO "mandated that hospitals poll each of their 6 patients at the end of their stay about whether their pain had been adequately treated."483 7 The scores that the hospitals received were 484 existentially important to the hospitals 671. 8 9 seeking to maintain their accreditation. "A low score puts a hospital in jeopardy of being ruled 10 ineligible for Medicaid reimbursements."485 11 672. 12 13 14 15 16 17 18 19 20 21 22 23 ⁴⁸¹ENDO-NY 00625759. ⁴⁸² Sari Horwitz, Scott Higham, Dalton Bennet and Meryl Kornfield, "Inside the opioid industry's marketing machine," 24 Post, December 2019. available https://www.washingtonpost.com/graphics/2019/investigations/opioid-marketing/ 25 ⁴⁸³ Melina Sherman, "Opiates for the masses: constructing a market for prescription (pain)killers," Journal of Cultural Economy, Vol. 10. 2017, available https://www.tandfonline.com/doi/full/10.1080/17530350.2017.1352010 26 ⁴⁸⁴ JCAHO has since abandoned the idea that pain is the fifth vital sign. 27 ⁴⁸⁵ Melina Sherman, "Opiates for the masses: constructing a market for prescription (pain)killers," Journal of Cultural Vol. 10. Issue 2017. available 28 https://www.tandfonline.com/doi/full/10.1080/17530350.2017.1352010 ⁴⁸⁶ ENDO-NY 00625759.

1 2 3 "Pain is now considered a fifth vital sign, along 4 with pulse, blood pressure, temperature and 5 respiration rate." 6 7 8 9 10 11 Source: Clinical (oxymorphone HCI) (ii) Management, April 2005 12 673. 13 14 15 16 17 18 487 19 The idea was simple: prescribe Opana, and you won't have to worry about low 674. 20 JCAHO scores. "It suffices to say that this system incentivizes the use of drugs – as reliable, fast-21 acting opioids for (temporarily) warding off the experience of pain. It is telling, after all, that the 22 23 uptake of these new systems and instruments of objectively measuring the subjective experience of 24 pain have paralleled steep increases in opioid prescriptions."488 25 26 27 ⁴⁸⁷ ENDO-NY-00625759

⁴⁸⁸ Melina Sherman, "Opiates for the masses: constructing a market for prescription (pain)killers," Journal of Cultural

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2017,

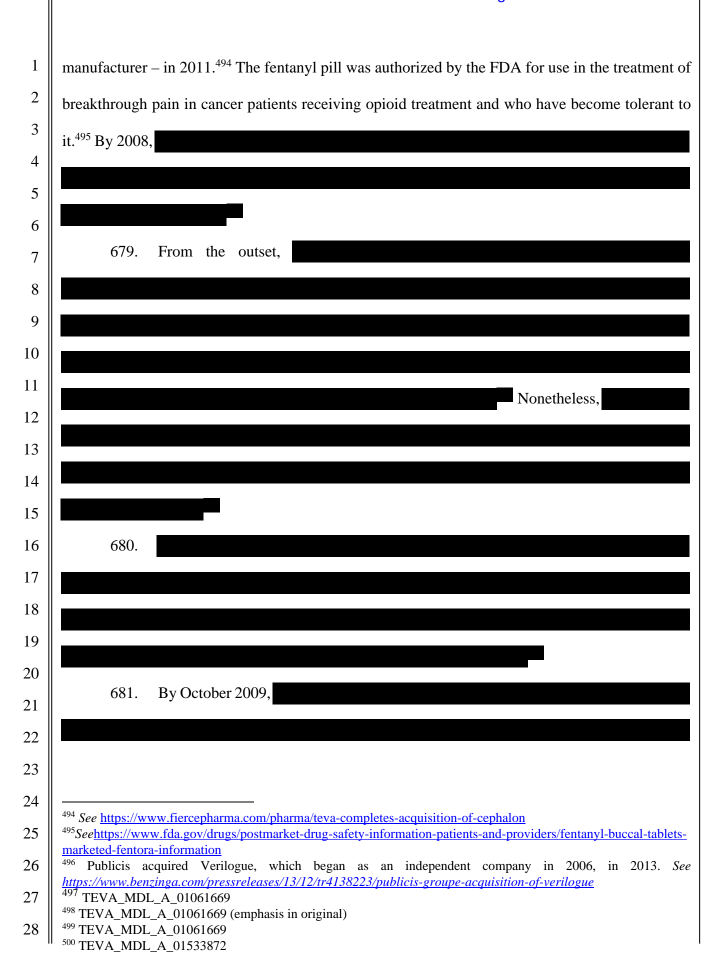
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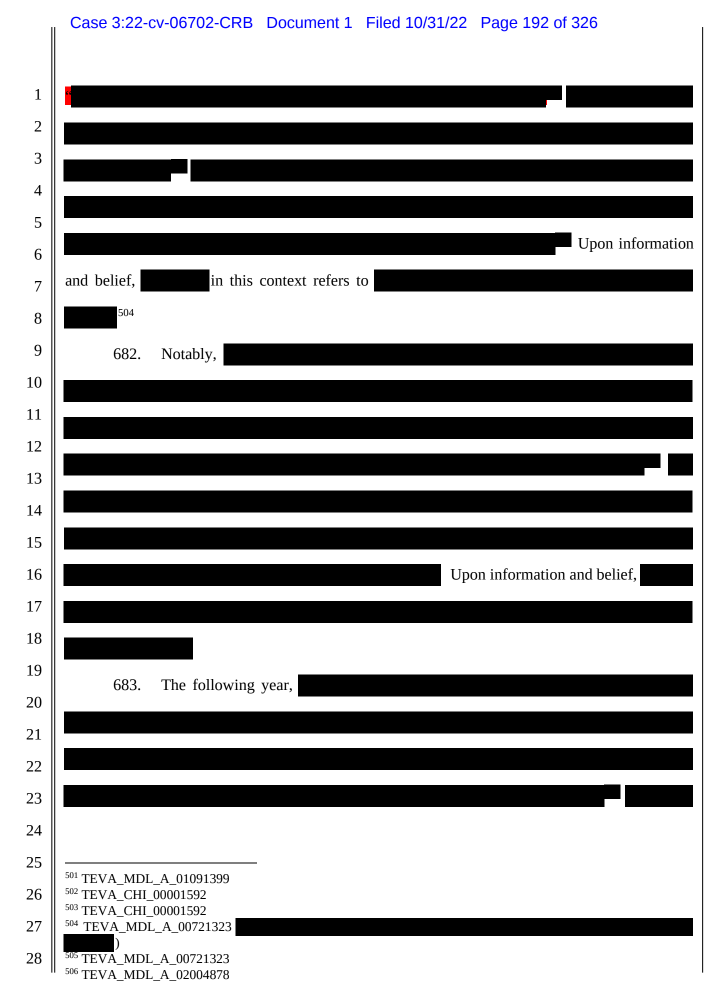
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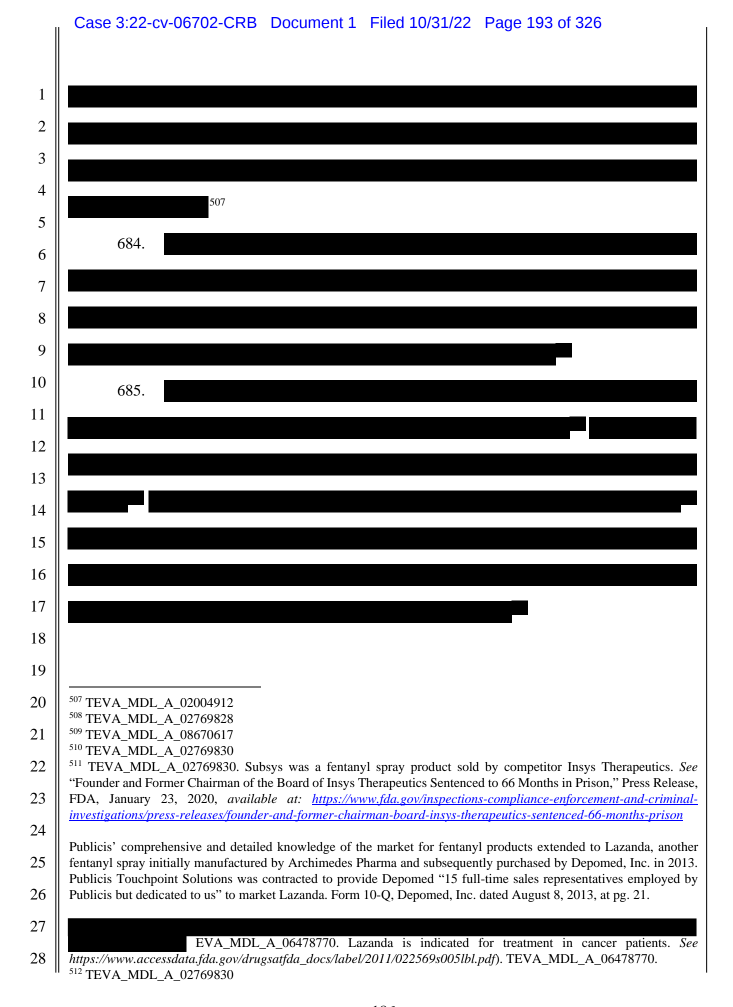
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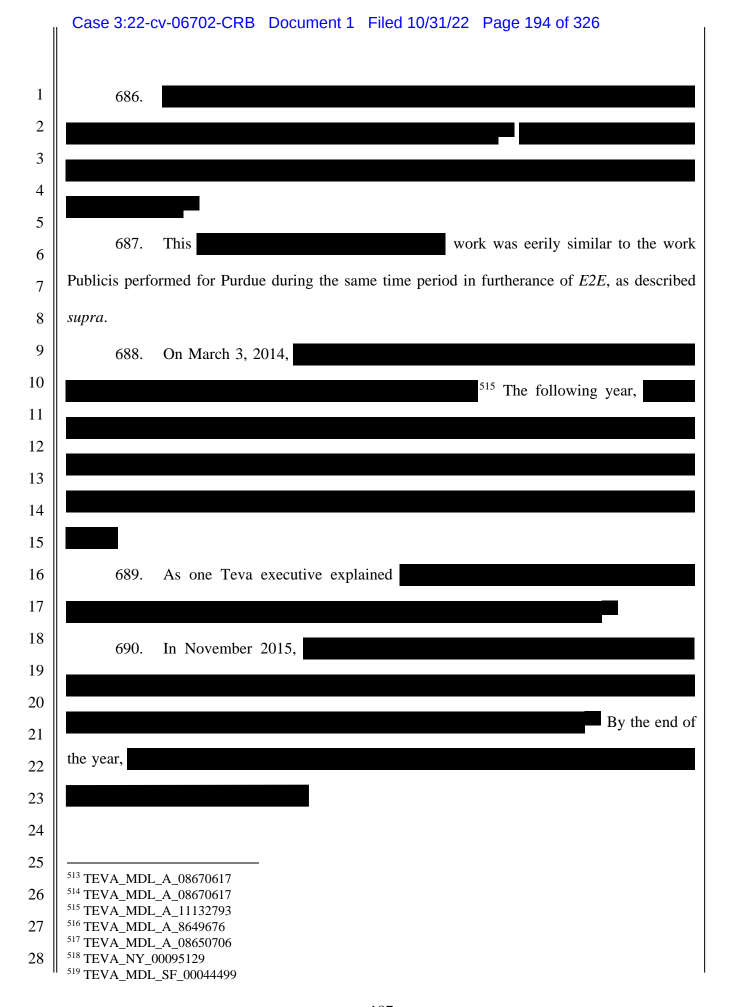
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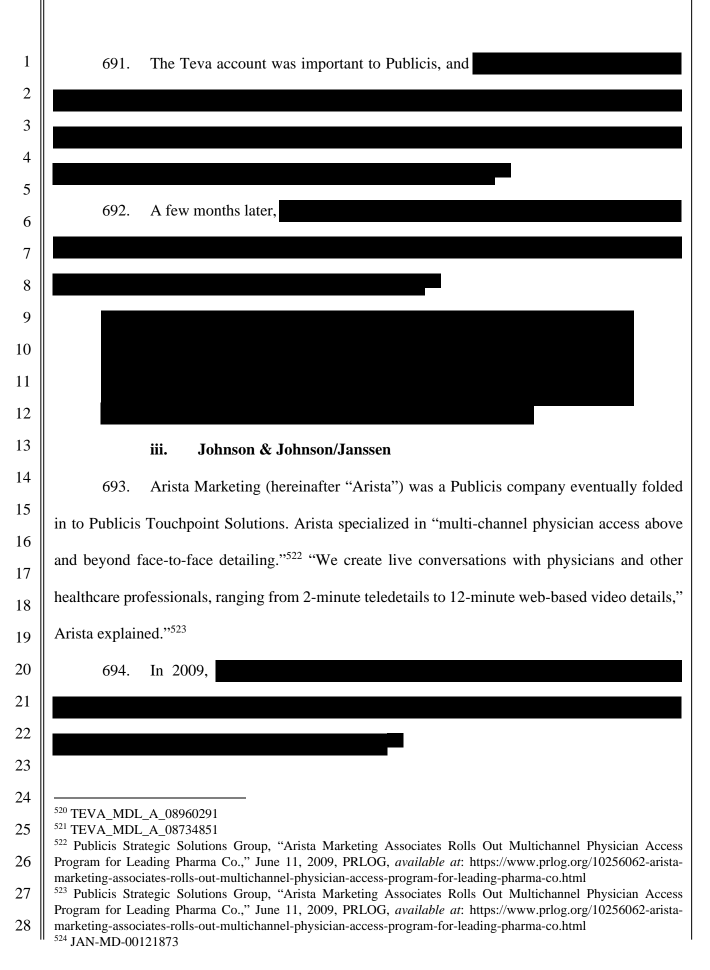
1 4. Beyond Saatchi: Endo's Broader Relationship with Publicis 2 675. Saatchi wasn't Publicis' only source of revenue from Endo. In January of 2012, 3 4 5 A few months later, in April of 2012, Endo 676. 6 7 8 9 10 11 12 13 14 Endo was keen on getting prescribers to switch from drugs like Vicodin to long 677. 15 16 acting opioids like Opana ER. 17 18 as well as 19 20 21 ii. Teva 22 678. Teva Pharmaceuticals ("Teva") also availed itself of Publicis' myriad services. Teva 23 24 sold Fentora, a fentanyl buccal tablet, after it acquired Cephalon, Inc. - Fentora's original 25 26 ⁴⁸⁹ ENDO-OPIOID_MDL-03744169. 27 ⁴⁹⁰ ENDO0111659 ⁴⁹¹ ENDO0111708 ⁴⁹² ENDO0111708 28 ⁴⁹³ ENDO0111708

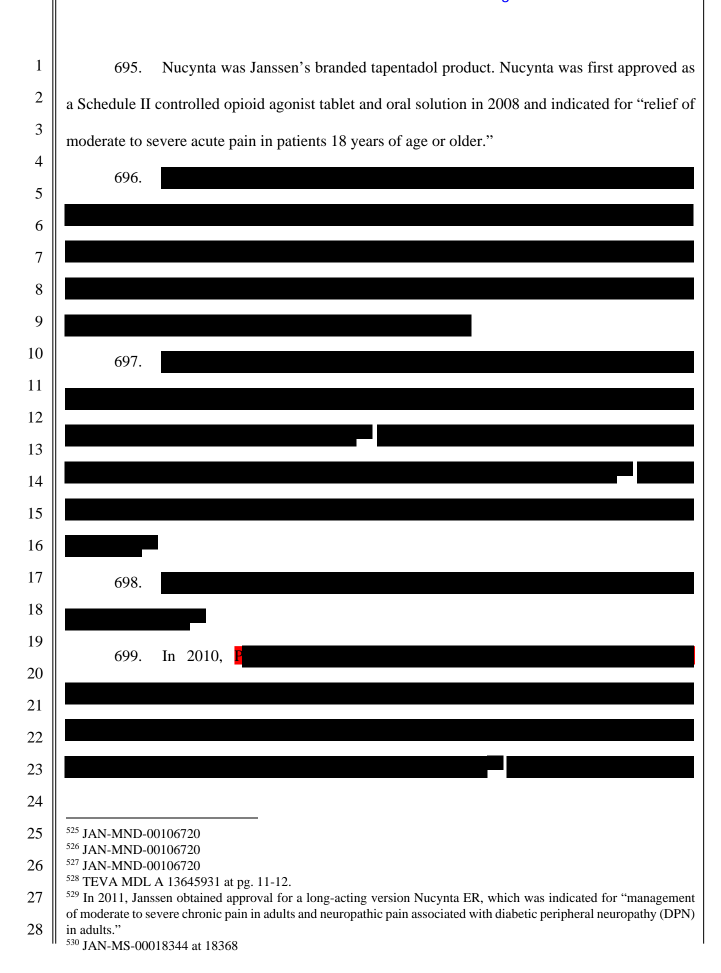












1 2 3 700. 4 5 6 7 8 9 701. Three years later, 10 Indeed, by 2013 11 while simultaneously working for competing products manufactured by 12 others. 13 **Servicing Manufacturers in Groups** iv. 14 702. Publicis did more than perform discreet work for individual opioid manufacturers; 15 16 it crafted industry-wide marketing efforts to boost sales not only of individual opioid products, but 17 of opioids *generally*. A rising tide lifts all boats. Or, to borrow the Purdue's executive's analogy, 18 Publicis worked to "make the pie bigger for all." 535 19 703. Accordingly, in addition to maintaining separate client relationships with multiple 20 opioid manufacturers, Publicis also worked for industry-wide groups to coordinate marketing and 21 advertising related to opioids, broadly. 22 23 24 25 531 JAN-MS-00018344 at 18368 26 ⁵³² JAN-MS-00772421 at 2, 5. ⁵³³ JAN-MS-00772421 at 6. 27 ⁵³⁴ JAN MS 01093081; JAN MS 01093079 535 See supra. at Paragraph 90; Sari Horwitz, Scott Higham, Dalton Bennet and Meryl Kornfield, "Inside the opioid 28 industry's marketing machine," 2019, available Washington December

https://www.washingtonpost.com/graphics/2019/investigations/opioid-marketing/

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704. Likewise,

v. Servicing Demand on the Comeback⁵³⁷ – Publicis and Orexo AB

705. The cognitive dissonance within Publicis as it embarked upon its work with Purdue on *E2E* must have been palpable. Three months after Publicis was working on OxyContin coupons, Swedish pharmaceutical company Orexo AB announced that the U.S. Food and Drug Administration had approved Zubsolv, its drug designed to treat opioid addiction. Zubsolv is a combination of buprenorphine and naloxone. At the time of Zubsolv's launch in 2013, Orexo projected peak sales of the drug to exceed \$500 million, annually.

706. But Orexo did not have a significant presence in North America when Zubsolv was approved. It did not have its own sales force in North America that could market its drug. As the most attractive market in which to sell opioid dependency treatments, Orexo desired to partner with someone who had the expertise and capacity to successfully launch Zubsolv in the United States.

707. On July 1, 2013, Orexo announced "that the company has entered into a commercial partnership with Publicis Touchpoint Solutions for the launch of Zubsolv in the United States...

⁵³⁶ JAN-NYDFS-0000123744

⁵³⁷ See Chris Rock, Bigger & Blacker, HBO TV Special, July 10, 1999, CR Enterprises, 3 Arts Entertainment, Production Partners, available at: https://www.youtube.com/watch?v=RRN3d58 MTk ("There ain't no money in the cure; the money's in the medicine. That's how you get paid... on the comeback. That's how a drug dealer makes his money. On the comeback.")

Publicis Touchpoint Solutions will be responsible for the execution of all field-based promotion activities through dedicated sales representatives and medical support to health care practitioners through deployment of a dedicated medical scientific liaison team."⁵³⁸

- 708. In announcing the partnership, Orexo Chief Executive Officer Nikolaj Sorensen emphasized Publicis' "knowledge of the opioid dependence therapeutic area," in addition to its expertise with "similar product launches," as primary reasons Orexo chose Publicis to be its partner.
- 709. As described in detail above, Publicis did indeed have expertise with similar product launches. While its sales representatives in Publicis Touchpoint Solutions were diligently selling Zubsolv for Orexo, Publicis worked just as diligently through its other agencies with other opioid manufacturers to maximize the sales of the drugs that were the direct source of Zubsolv's indication. The same year it partnered with Orexo to launch Zubsolv, Publicis billed Purdue Pharma around \$8mm for marketing work on OxyContin and other Purdue opioid products.
- 710. Most incredibly, Publicis began work on Purdue and McKinsey's *E2E* project **less than three months** after its partnership with Orexo was announced.
- 711. Through its work with Orexo, Publicis gained knowledge of the market opportunities created by widespread opioid dependency, and of the co-dependence of the markets for opioid treatments and opioid use disorder treatments.
- 712. As Orexo, Publicis' partner, described in its 2013 annual report, "[p]rescription painkillers containing opioids are highly addictive, and regular or long-term use can lead to physical dependence." Orexo further observed that "[m]any abusers begin by taking opioids orally," and that the misuse of opioid prescription drugs was a "growing problem," with "opioid dependence more common than the abuse or, or dependence on, any other type of prescription medication."

⁵³⁸ "Orexo Forms a Commercial Partnership with Publicis Touchpoint Solutions for Launch of Zubsolv in the US," July 1, 2013, *available at:* https://www.businesswire.com/news/home/20130701005515/en/Orexo-Forms-a-Commercial-Partnership-with-Publicis-Touchpoint-Solutions-for-launch-of-ZubsolvTM-in-the-US

713. Describing the addressable market for Zubsolv, Orexo stated that the "cost of prescription opioid abuse, dependence and misuses in the US is estimated to exceed USD 56 billion per year," and further, "15,000 people die from opioid pain relievers each year in the US. Deaths from opioid pain relievers exceed those from all drugs and traffic accidents."

714. The conclusion was obvious:

Zubsolv has entered a large and growing market. The current US market of products containing buprenorphine/naloxone amounts to approximately USD 1.9 billion, before rebates to payers, co-ay support and other discounts. The market continued to grow by 9 percent in value and 10 percent in volume during 2013. Continued double-digit growth is likely in the years to come, and will be driven by the significant unmet medical need, the growing number of opioid dependent patients as well as the impact of the Affordable Care Act. 539

- 715. Publicis Touchpoint Solutions provided a contract sales force to Orexo through the second quarter of 2014, and continued to advise its client on Zubsolv sales through at least 2019.⁵⁴⁰
- 716. By 2015, Orexo stated flatly, "[t]here is no doubt opioid addiction has become one of the major health concerns in the US and a disease currently out of control." The market for Zubsolv was booming, with an 83% increase in revenue compared to the prior year. 542
- 717. Throughout its time selling a treatment for opioid use disorder, Publicis endeavored on multiple fronts, with multiple clients, to maximize the sales of opioids, the drugs that cause the condition that Zubsolv another drug it was paid to sell treats. The synergies were hard to beat. Publicis helped Orexo service a demand it had a principal role in creating and sustaining through its *contemporaneous* work on *E2E* and other projects. Indeed, without Publicis efforts promoting opioids for its other clients, there may not have been an addressable market for Orexo's product in the first place.

n. Alternative Channels: Publicis and Practice Fusion

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⁵³⁹ See http://mb.cision.com/Main/694/9557375/224609.pdf (emphasis added).

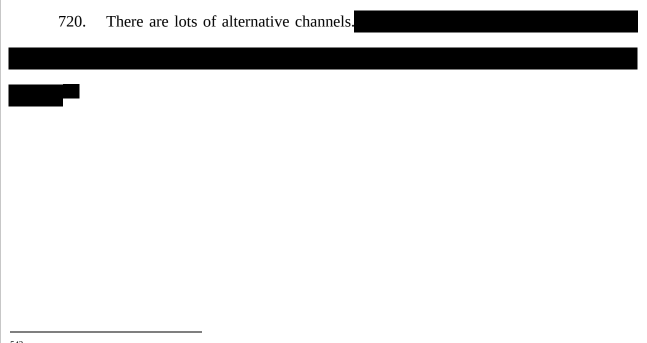
⁵⁴⁰ See https://www.linkedin.com/in/susan-broadnix-8794bb7/

⁵⁴¹ See https://mb.cision.com/Main/694/9942119/491960.pdf

⁵⁴² *Id*

718. In pharmaceutical sales, the traditional workhorse of sales and marketing campaigns is the individual sales representative calling on individual prescribers to meet in-person with them in order to extol the virtues of a given drug and encourage the doctor to prescribe more of it to patients. Classically, this has been the primary channel through which a pharmaceutical company's message has been delivered to the intended audience.

719. Of course, there are alternatives to face-to-face meetings with doctors by sales representatives. In theory, wherever a prescriber's attention is focused at any given moment is a potential spot to deliver content to her. Over the past few decades, for instance, doctors increasingly interact with screens that they look at while at work. Forty years ago, doctors didn't walk around their offices holding iPads, but they do now. This is an "alternative channel" through which a message can be delivered, and Publicis endeavored with its clients to create multi-faceted campaigns so that a prescriber would be surrounded, in effect, with pharmaceutical company messaging coming from all directions: the people he has lunch with, the speakers at conferences he attends, the ads on his screen, the search results on his computer when he uses Google to search for a condition state, etc.



⁵⁴³ PPLPC014000224450.

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Channel	Potential Targets	Potential Tactics	Vendors	Cost	Impact
Associations	Targets who are members	Approved messaging; educational focus	APS, AAPM, AAFP, ACP, AAOFP, AAPA, AAPM&R	Med	Med
EHR	EHR-using targets	POC messaging	Practice Fusion	High	Med
			AllScripts		
In-office	 High-value targets due to cost 	Customized wall charts	McCallan Health	High	Med
			Accent Health		
			Media Health Network		
Surveys	All targets	ATU-style questions	WorldOne Interactive	Low	Med
Nurse Focus	All NP targets	Same messaging as MD	RNSights	Med	High
Pharmacist Focus	All Pharmacy targets	ADF/access messaging	SK&A	Med	Med
MC Focus	All MC targets	ADF/access messaging	SK&A	Med	Med
Contract Field Force	• IDNs	Standard rep messaging	Best MSLs	High	Med
Social	All targets	Educational messagings	Doximity	Med	Med
			QuantiaMD		
KOL Programs	All targets	KOL on-demand	Synapse	High	Med
Conference Services	Attendee lists	High-value messaging	Pain-related conferences	Med	High
Authenticated Website	- All targets	High-value messaging	UBM Medica	High	Med
eSample	Sample-appropriate targets	Sampling	Physicians Interactive	High	Med
	(Intermezzo)		Doctor Directory		
Website	All targets	All messaging	PurdueHCP.com	Med	High

721. These channels, when utilized together as part of an overall package, can be said to comprise the overall "marketing mix" that Publicis, ZS, McKinsey and others continually endeavor to optimize and espouse as a "best practice," which is another way of saying that lots of the consultants' other clients do the same thing.

Publicis stood athwart the industry, in a position to advise their clients on which vendors to pick to best utilize these alternative channels. ⁵⁴⁴ In this way, as in others, it served as a hub, or an intermediary connecting multiple participants in the overall efforts to market opioids to maximum effect.

722. "EHR" stands for Electronic Health Record. It is a relatively new marketing channel, and one whose growth was spurned by federal legislation in 2009 that encouraged doctors to adopt electronic record-keeping methods for patient records. As early as 2012, Publicis was working with Purdue to develop banner advertising and other types of content that could be placed in the EHR

 $^{^{544}}$ Indeed, this position athwart the distribution channels is what being an AOR is.

channel. This work continued at late as 2018, and Publicis, as AOR, also selected EHR vendors in which the content could be placed.

- 723. They chose Practice Fusion.
- 724. As the varieties of marketing channels above make clear, pharmaceutical marketing is complex. Publicis Health's Chief Digital Officer recently remarked on his complexity:

When we think about our ecosystem, the complexity goes beyond just regulatory. We are in a complex ecosystem in the sense that we are marketing to two different audiences in a patient, and a physician. We're hoping that they come together and have a productive conversation, and in that conversation, we're hoping that they talk about the treatment that we're marketing to them. And that's complex, I don't think there's any other industry that is met with that challenge.⁵⁴⁵

725. This complexity creates reams of data. ZS Associates, the pharmaceutical sales and marketing consultancy, has described the "endemic challenge" facing the pharma industry: "It has a complex and circuitous sales process involving drug manufacturers, physicians, pharmacies, patients, and insurance companies. Each step in a buying process creates data – and more of it is being created every day."⁵⁴⁶

i. Electronic Health Records – A New Frontier

726. These data streams may be used as a tool. The data available and the information provided by EHR is all-encompassing in its breadth. As ZS Associates noted, "[t]he spectrum of data is almost unprecedented: lab results, diagnoses, prescriptions, patient compliance, physician notes, and follow-on or replacement prescriptions."⁵⁴⁷ The result is that EHR data can provide an

Ray Rosti, Chief Digital Officer of Publicis Health Media, December 7, 2020, available at https://www.youtube.com/watch?v=AG4HLIDSQ A

⁵⁴⁶ Steve Love, Sudhanshu Bhatnagar, Greg Rickman, and Jedy Wang, "The Value of EMR Data: Unlocking Insights That Drive Pharma Sales," *Journal of Pharmaceutical Management Science Association*, Spring 2016, *available at*: https://llibrary.net/document/zlewp5lq-value-emr-data-unlocking-insights-drive-pharma-sales.html .

In 2011, ZS touted Publicis' endorsement of its own product offerings, such as Javelin. *See* https://twitter.com/ZSAssociates/status/119892005756215297

⁵⁴⁷ Steve Love, Sudhanshu Bhatnagar, Greg Rickman, and Jedy Wang, "The Value of EMR Data: Unlocking Insights That Drive Pharma Sales," *Journal of Pharmaceutical Management Science Association*, Spring 2016, *available at*: https://llibrary.net/document/zlewp5lq-value-emr-data-unlocking-insights-drive-pharma-sales.html.

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end-to-end perspective for the entirety of a patient's journey from the diagnosis of a disease through treatment and through cure or death.⁵⁴⁸

727. This data can be used to sell more drugs. With the proliferation of electronic health data, vendors have arisen to aggregate and sell data sets and analytics platforms based on this data. These providers "now give pharma companies greater visibility than ever into its (*sic*) marketplace of buyers, consumers, and decision-makers – and the factors that drive sales."⁵⁴⁹

ii. What Practice Fusion does

728. Practice Fusion was founded in 2005 as a vendor of free electronic health record storage (EHR) software. It "makes software that many doctors see on their devices. When you go into the exam room, your electronic records pop up on their screens." 550

729. Practice Fusion's cloud-based EHR software platform serves tens of thousands of active health care providers in the United States, and the software is used during millions of physician-patient encounters each month. It is "the #1 cloud-based ambulatory EHR platform in the U.S., supporting 30,000 medical practices in delivering better care to 5 million patients a month. With a best-in-class satisfaction rate, Practice Fusion is committed to delivering intuitive and easy-to-use health IT solutions to small, independent medical practices." 553

730. The business model was daring; most EHR vendors – Practice Fusion's competitors
 – charge users to use their software platforms, typically in the form of a licensing fee. Practice

⁵⁴⁸ *Id*.

⁵⁴⁹ *Id*.

⁵⁵⁰ Brian Mann, "Health Care Software Firm Fined \$145M in Opioid Scheme With Drug Companies," *NPR*, February 1, 2020, *available at:* https://www.npr.org/2020/02/01/801832788/healthcare-software-firm-fined-145m-in-opioid-scheme-with-drug-companies

⁵⁵¹ SK&A, Report on Physician Office Usage of Electronic Healthcare Records Software (February 2016).

⁵⁵² Reaction Data, Report on EHR Satisfaction According To Physicians (January 2018).

⁵⁵³ Practice Fusion Company Profile, https://www.practicefusion.com/about/

1 Fusion was different. Ryan Howard, founder and CEO of Practice Fusion, explained, "our product 2 being free and web-based was incredibly unorthodox."554 3 731. The company didn't charge the doctors for these intuitive and easy-to-use health IT 4 solutions. It was free to them.⁵⁵⁵ Instead, Practice derived revenue from payments from 5 pharmaceutical companies in exchange for ad space, and other marketing products like "clinical 6 decisions alerts" (CDS) in its EHR software that served as advertisements for the pharmaceutical 7 company's products.⁵⁵⁶ 8 9 Practice Fusion marketed its platform to the pharmaceutical companies as a way to 732. 10 influence prescriber behavior. For instance, Practice Fusion's pitch materials to pharmaceutical 11 companies indicated that a pain CDS could be aligned with that company's "brand objectives." 12 733. From the advertiser's perspective, this product offering was tantalizing. 13 14 15 16 17 18 "Practice Fusion, the cloud-based 19 electronic medical system, mirrors the real-life workflow of a doctor's office," explained John 20 Mehta, Publicis Sapient's Chief Experience Officer from 2019 to 2020. 559 21 22 554 Ryan Howard, CEO Practice Fusion, Cracking the Entrepreneur Code, September 26, 2011, available at: 23 https://www.youtube.com/watch?v=JaTIgMUEc9Y&t=1067s 555 "If something is free, you're the product." See Richard Serra, Carlota Fay Schoolman, Television Delivers People, 24 1973 ("The Product of Television, Commercial Television, is the Audience... Television delivers people to an advertiser."), available at: https://www.youtube.com/watch?v=LvZYwaQlJsg 556 Christina Farr, "Practice Fusion, once headed for \$1.5 billion valuation, ends in 'disappointing' fire sale," CNBC, 25 January 8, 2018, available at: https://www.cnbc.com/2018/01/08/practice-fusion-acquired-by-allscripts-for-100-26 million-in-cash.html (""Rather than selling expensive licenses, the company makes money by showing ads to physicians that use the service.") ⁵⁵⁷ See 27 TEVA CHI 00437579. TEVA CHI 00437579. 28 ⁵⁵⁹ John Maeda, "Better Health by Design: Making Healthcare Tech More Usable, Understandable And Profitable, TechCrunch, December 8, 2015, available at: https://techcrunch.com/2015/12/08/better-health-by-design-making734. The brand objectives that Practice Fusion pursued for Purdue and other opioid manufacturers included expanding the market share of a given company's extended-release opioids as well as growing the *overall* size of the ERO opioid market by targeting "opioid naïve" patients and patients that were on immediate release ("IRO") opioid therapies.

735. Practice Fusion's EHR platform provided Purdue and other opioid manufacturers a way to reach into the examination room and interact with the prescriber and the patient by utilizing the "wealth of data" available in EHR records⁵⁶⁰ to deliver targeted messaging when and *when* it matters. Moreover, Practice Fusion provided its customers (pharmaceutical companies, not doctors) with "novel tools" to "drive appropriate care for patients with chronic pain," by utilizing private, individualized patient health care records in order to target the delivery of messages intended to increase overall ERO prescribing. The platform enabled Practice Fusion's customers to insert promotional messaging throughout the provider workflow, including during patient visits and "patient-centric provider targeting." ⁵⁶¹ Practice Fusion provided these services to its customers so without patient or physician consent. ⁵⁶²

736. As early as 2012, Publicis was working with Purdue to develop banner advertising and other types of content, including CDS alerts, that could be placed within the EHR channel. This work continued at late as 2018, and Publicis, as AOR, also selected EHR vendors in which the content could be placed: Practice Fusion.

also

⁽identifying Maehda as Publicis' Chief Experience Officer. According to LinkedIn, Maehda left Publicis Sapient in October of 2020. *See* https://www.linkedin.com/in/johnmaeda/).

⁵⁶⁰ See PFDPA00000025-27 (noting the wealth of data gleaned from access to the medical records helped build a franchise to treat patients in pain "around the clock" and expand the data sets of conditions for which to prescribe opioids).

⁵⁶¹ *Id*. at 4.

⁵⁶² Press Release, Department of Justice, Electronic Health Records Vendor to Pay \$145 Million to Resolve Criminal and Civil Investigations, Practice Fusion Inc. Admits to Kickback Scheme Aimed at Increasing Opioid Prescriptions (Jan. 27, 2020), https://www.justice.gov/opa/pr/electronic-health-records-vendor-pay-145-million-resolve-criminal-and-civil-investigations-0

iii. Publicis and Practice Fusion

737. Publicis piloted Practice Fusion to Purdue over years of working with both. In one sense, this is Publicis performing its role as AOR for Purdue. The AOR not only *creates* the content, but *places* it as well. The AOR acts as a distributor of the client's ads to people selling space in which to put them. A billboard on the side of an interstate, a prospective patient's TV screen, signage on the wall above the urinals in a barroom bathroom, a prescribers' iPad screen... it's all placement. Choosing the best places and putting Purdue's messages in them was part of Publicis' job as AOR.

738. One of the best spots, Publicis informed Purdue, was Practice Fusion. In the simplest of terms, Practice Fusion had advertising space to sell in the form of banner advertising and embedded features such as CDS alerts that could be placed within Practice Fusion's software programs that prescribers looked at on their screens throughout the day, and Publicis put some Purdue content there.

739. In October 2013,

563 Practice Fusion's subsequent work with Purdue was carried out in coordination and conjunction with Publicis, and as a part of the McKinsey's

implementation of E2E alongside Purdue to

- 740. Publicis even paid Practice Fusion's invoices on behalf of Purdue.
- 741. Publicis and Practice Fusion coordinated their work for Purdue. In April 2014, with *E2E* in full swing, Publicis' John Dwyer reached out to Practice Fusion's Jim Pantaleo to discuss some "aggressive EMR strategies" for Purdue, and focused on the point of prescription, when the

PPLPC018000873870.

⁵⁶³ PPLPC014000224450; PPLPC014000224448 (

an immediate-release opioid (IRO), an ad featuring Oxycontin's 12-hour dosing would appear. A doctor contemplating an extended-release opioid (ERO) would see an ad touting OxyContin's formulary coverage. Dwyer also suggested a "reassessment" reminder appear within the EMR each time a patient receives a consecutive prescription for OxyContin, in which the prescriber would be reminded of Purdue's titration messaging, described above. Dwyer suggested four such "aggressive" strategies, and suggested that the strategies that have the "highest ROI" and are the best "Rx drivers" be reinvested in:

doctor is deciding whether and what to prescribe to a patient. For doctors contemplating prescribing

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To: John Dwyer[john.dwyer@rosetta.com]

From: Jim Pantaleo

Sent: Mon 8/11/2014 9:49:51 AM (UTC-04:00)

Subject: RE: Purdue Brands - EMR Strategy/Ideas/Discussion

Thanks John. Checking in sounds like a good plan.

In the meantime, I will put together messaging examples of how others are targeting in the platform and send to you.

We may not have the exact strategic messaging tactics, but I believe all of the below can be executed.

I'll send by end of day; first thing tomorrow.

From: John Dwyer [mailto:john.dwyer@rosetta.com]

Sent: Monday, August 11, 2014 9:36 AM

To: Jim Pantaleo

Subject: Re: Purdue Brands - EMR Strategy/Ideas/Discussion

Hi Jim,

I agree, and I think we should try to check in with each other for a few minutes every other week or so while these programs are of high interest to Purdue.

Here are the 4 approaches we discussed on Friday's meeting. Again, with these approaches, the most helpful thing you can provide would be examples of other products that have used a targeting strategy to serve ads based on competitor brands or generics in the same category. EMRs for EROs (addresses Conversion, Titration, opioids with abuse-deterrent properties (OADPs), Mgd Care)

· Pilot 3 aggressive EMR strategies in Q1 and quickly optimize and reinvest in the 1-2 highest ROI and Rx drivers

- 1. Target IRO oxycodone Rxs with branded OxyContin Q12h msg or unbranded ER teaser msg
- 2. Serve a "Reassessment" reminder msg each time pt receives consecutive OxyContin Rx
- 3. Target non-ADP Rxs with branded OxyContin ADP msg or unbranded ADP teaser msg
- 4. Target ERO Rxs with OxyContin MC coverage message or unbranded coverage teaser msg

Thanks again for coming by on Friday and sharing so much information.

John Dwyer

Associate Partner, Healthcare O: 212 771-5109 M: 917 797-9317

99 Hudson St. 7th Fl New York, NY 10013 USA

Rosetta.com

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742. Three months later, in July 2014, Publicis proposed that EHR channels be used to

encourage conversion of immediate-release opioids (IRO) patients to extended-release opioids 26

(EROs). This would have the effect of growing the overall size of the ERO market by creating new

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ERO users who previously were prescribed IROs. One idea was to target any patient who has been prescribed an IRO "multiple times (3 or 4) with a branded or unbranded message for OXC."

743. Consistent with the overall goals of *E2E*, Publicis encouraged Practice Fusion to help grow the ERO market by encouraging conversions of IRO users. Publicis informed Purdue that Practice Fusion had the relevant capabilities: the ability to deliver "Branded OxyContin & clinical messages... during patient eRx, patient Medication Selection... and edit patient Medication." Publicis' John Dwyer asked his colleagues whether Practice Fusion had "any kind of ROI or Rx impact metrics around these that they can share?"

iv. Practice Fusion and Purdue

- 744. In April of 2013, Publicis agreed to Purdue place banner advertisements inside of Practice Fusion's EMR software. The banner ads would appear when a prescriber was seeing a patient with certain health conditions, and the Purdue ads would appear with the patient's EHR suggested pain-related content (for instance, a reference to the patient's "pain symptoms."). Three of the banner ads were for OxyContin, two were for Butrans.
- 745. After a few months, because of compliance concerns, Purdue paused the banner advertisement initiative, but in December 2013, Publicis made the case for continuing the initiative, and told Purdue, "Practice Fusion is the only partner that offers Banners within workflow prior to making a prescribing decision. Helps to increase awareness and ultimately sales." 565
- 746. Meanwhile, Practice Fusion continued to solicit business from Purdue directly. In May 2014, for instance, Practice Fusion sent a news article to Purdue about Practice Fusion's implementation of a CDS program for a vaccine manufacturer. The article was forwarded internally at Purdue to the new CEO, Mark Timney, who had been appointed CEO four months prior. Timney responded, "Thanks. The key is understanding how it grows or protects scripts." 566

⁵⁶⁵ Massachusetts Publicis Complaint Para. 103.

⁵⁶⁶ Practice Fusion Information at Para. 28.

v. CDS Alerts: Circumventing No-Sees, Influencing Prescribing, Making Money

747. What is a CDS Alert? A doctor using Practice Fusion's software would see a message "alerting the healthcare provider that, given the particular personal health information and circumstances of the patient before the provider at that moment, the provider should consider certain clinical information, perform certain tests or assessments, and complete certain documentation." ⁵⁶⁷

748. Do CDS alerts work?⁵⁶⁸ At a 2016 seminar for pharmaceutical companies put on by Ogilvy (another advertising agency, owned by Publicis competitor WPP), Practice Fusion presented, and gave the audience an example of the of power of the clinical decision support program:⁵⁶⁹

Practice Fusion, a cloud-based EHR provider, offered an example in which it launched an obesity clinical decision support program. The HER notified physicians with messages at the point of care about recording patients' BMI stats and, if high, noting a treatment plan. The program reached more than 50,000 physicians and 3.7 million patients resulting in 25,000+ more patient plans, which was a 5-fold increase.

749. Why do CDS alerts work? Well, "the closed system of EHR's... means the marketing and communications from pharma to physicians are not scattershot web ads, but much more targeted inside communications that can inform doctors at a critical moment." Presenter

⁵⁶⁷ Practice Fusion Information at Para. 17.

⁵⁶⁸ "Work," in the sense of making money for the pharmaceutical company. ROI is the relevant metric. If the product cannot increase sales for the pharmaceutical company that pays Practice Fusion, no pharmaceutical company will pay Practice Fusion money, and the company would go broke or have to "pivot" to another busines model and pray for rain. Practice Fusion was not a non-profit, nor a B Corp. It was in business to make money.

⁵⁶⁹ Beth Snyder Bulik, "Ogilvy CommonHealth takes deep dive on EHR as a marketing strategy for pharma clients," *Fierce Pharma*, May 9, *available at*: https://www.fiercepharma.com/marketing/ogilvycommonhealth-takes-deep-dive-ehr-as-a-marketing-strategy-for-pharma-clients (emphasis added).

⁵⁷⁰ Beth Snyder Bulik, "Ogilvy CommonHealth takes deep dive on EHR as a marketing strategy for pharma clients," *Fierce Pharma*, May 9, *available at*: https://www.fiercepharma.com/marketing/ogilvycommonhealth-takes-deep-dive-ehr-as-a-marketing-strategy-for-pharma-clients (emphasis added).

after presenter noted not only the value of EHRs but also that using them needs to be thought of as a strategy, not just a tactic or channel."⁵⁷¹

- 750. Could CDS alerts be used as a strategy (or tactic) to maximize sales of a controlled substance that the planners know is causing widespread addiction, abuse, and death? Yes.
 - 751. Were they? They were.
- 752. In late 2015, Purdue gave its approval to move forward with the Pain CDS alert program. Because the overall goal of the program was to increase the size of the *overall* ERO market, each of Purdue's three ERO brands split the cost of the Practice Fusion program.
- 753. By 2016, Purdue and Practice Fusion were working to use CDS alerts for the "objective" to "Grow ERO prescriptions within the Practice Fusion eHR." Publicis estimated that the return on investment (ROI) on the CDS alerts would be 2:1.⁵⁷² An additional \$2 million in annual revenue was possible by using Practice Fusion to convert IRO patients to ERO patients, but it would cost \$1 million annually in payments to Practice Fusion in order to obtain that additional revenue:

⁵⁷¹ Beth Snyder Bulik, "Ogilvy CommonHealth takes deep dive on EHR as a marketing strategy for pharma clients," *Fierce Pharma*, May 9, *available at*: https://www.fiercepharma.com/marketing/ogilvycommonhealth-takes-deep-dive-ehr-as-a-marketing-strategy-for-pharma-clients (emphasis added).

⁵⁷² Practice Fusion performed its own ROI analysis of the Pain CDS program for Purdue, and estimated that Purdue would enjoy a decidedly rosier 5.8 to 7.8 times its cost. Practice Fusion Information at Para. 37. Just like Publicis, Practice Fusion was at all times focused on delivering ROI for Purdue. In an April 22, 2015, internal email, a Practice Fusion employee conceded, "Since this is being sent to a marketing audience the idea of ROI has to be part of the plan to justify the costs of the program." Practice Fusion Information at Para. 42.

Notably, this ROI analysis was left *out* of the written presentations that Practice Fusion provided to Purdue, as it would raise compliance concerns. "Don't include the ROI in the [written] proposal. We'll walk the client the ROI," advised the same employee who stated that "the ROI has to be part of the plan." Practice Fusion Information at Para. 43. Instead, Practice Fusion "voiced over" the "commercial impact" of its proposed program, instead of creating a paper trail.

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Estimated ROI Patients data captured within Practice Fusion eHR: Number of Patients with Chronic Pain taking IRO 1,100,000 Number with average Pain Score of 5+ 150,000 % Switched as a result of Quality Score Initiative 15% Patients switched to ERO 22,500 Purdue Share of switches 25% Average value of Switch in Purdue Revenue \$350 Revenue Generation \$2,000,000 Investment \$1,000,000 ROI 2:1 Butrans

- 754. Practice Fusion and Purdue entered into a Statement of Work on March 1, 2016, to provide a CDS program "directed at chronic pain management treatment with immediate release opioids and chronically used NSAIDs." ⁵⁷³
- 755. The contract stated that the "Parties agree and acknowledge that the collaboration project will follow national evidence-based guidelines, and will not encourage the prescribing or utilization of Purdue-specific product or services." Both Practice Fusion and Purdue knew at the time of contract formation this mutual representation was false. In fact, "national evidence-based guidelines" such as the CDC Guidelines released on March 15, 2016, were known to the Parties while the CDS was designed and implemented, but those guidelines were ignored. Likewise, a 2016 New England Journal of Medicine article entitled "Opioid Abuse in Chronic Pain Misconceptions and Mitigation Strategies counseled against the use of opioids for chronic pain, where the benefits of opioids were "much more questionable" than in the acute treatment context. Purdue and Practice

⁵⁷³ Practice Fusion Information at Para. 75.

⁵⁷⁴ Practice Fusion Information at Para. 78.

Fusion reviewed the article but ignored its conclusions. Instead, they designed the CDS alerts to *convert* users of IROs to EROs for long-term maintenance of chronic pain.

756. Purdue's marketing team – not Purdue's medical experts – worked with Practice Fusion to *design* the CDS and determine its functionalities. For instance, the CDS was designed to incorporate patient's Pain Score and a brief pain inventory ("BPI"), and Purdue's marketing staff also contributed to the design of the Care Plan options presented within the CDS, and the logic of the CDS software functionality itself.⁵⁷⁵ "BPI can increase ERO use," Purdue noted.⁵⁷⁶

757. The CDS program was launched on the Practice Fusion platform in early July 2016. The final pain CDS contained three separate alerts: (1) the first alert urged the healthcare provider to record a pain score; (2) the second alert recommended that healthcare providers take a brief pain inventory ("BPI") of patients that met a certain threshold, for patients who had a chronic pain diagnosis, and for patient who recorded two or more pain scores of four or more in the previous three months (utilizing a zero to ten point scale); (3) the third alert suggested the creation of a follow up plan to treat the patient's pain, which alert appeared when a patient reported a pain scale of four or higher within four months, or if a patient with chronic pain had a BPI contemplated.

758. The pain CDS alert implemented by Practice Fusion deviated from established medical guidelines by directing providers to record a treatment plan only when pain was classified as chronic or was above a certain threshold over a period of time. It did not incorporate the substance of the New England Journal of Medicine article from which the CDS sourced a list of treatment options.

759. Moreover, the Clinical Quality Measures ("CQM") performance standards require providers to record a treatment plan any time the pain assessment was documented as positive. Contrary to accepted medical practice, the pain CDS alert listed EROs as a treatment option on

⁵⁷⁵ Practice Fusion Information at Para. 90.

⁵⁷⁶ Practice Fusion Information at Para. 91.

equal footing with IROs and non-opioid therapy. Likewise, it also listed EROs as a treatment option for opioid naïve patients without regard to whether the provider had the adequate expertise to prescribe EROs.

- 760. As the CDS alerts went live, Practice Fusion kept Purdue updated on progress. Practice Fusion told Purdue that through November 30, 2016, the pain CDS alert had produced alerts during 21 million patient visits, involving 7.5 million patients and 97,000 healthcare providers.
- 761. Practice Fusion further indicated that since its Pain CDS alerts went into effect "there is a general shift toward EROs from IROs," the biggest impact recorded "within Emergency Medicine, Orthopedics, and Pain Medicine."
- 762. The Pain CDS alert was live on the Practice Fusion platform from early July 2016 to the spring of 2019. The Pain CDS alerted more than 230,000,000 times during this period. Physicians wrote hundreds of thousands of ERO prescriptions after one of the Pain CDS alerts had been triggered. Moreover, healthcare providers who received Practice Fusion's Pain CDS alerts prescribed EROs at a higher rate than those that did not.⁵⁷⁷

vi. Guilty

763. In January 2020, Practice Fusion paid \$145 million and entered into a deferred prosecution agreement with the Department of Justice for these CDS alerts and its work with Purdue.⁵⁷⁸ Nine months later, Purdue pled guilty for conspiring with Practice Fusion to violate the Anti-Kickback Statute by paying Practice Fusion for the CDS alerts which were intended to

⁵⁷⁷ See Practice Fusion Information, at Paragraph 115.

⁵⁷⁸ Press Release, Department of Justice, "Electronic Health Records Vendor to Pay \$145 Million to Resolve Criminal and Civil Investigations, Practice Fusion Inc. Admits to Kickback Scheme Aimed at Increasing Opioid Prescriptions", Department of Justice (Jan. 27, 2020), *available at*: https://www.justice.gov/opa/pr/electronic-health-records-vendor-pay-145-million-resolve-criminal-and-civil-investigations-0">https://www.justice.gov/opa/pr/electronic-health-records-vendor-pay-145-million-resolve-criminal-and-civil-investigations-0 (describing Practice Fusion's conduct as "abhorrent" and noting Practice Fusion and Purdue "illegally conspired to allow [Purdue] to have its thumb on the scale at precisely the moment a doctor was making an incredibly intimate, personal, and important decisions about a patient's medical care, including the need for pain medication and prescription amounts.")

increase sales of Purdue's drugs.⁵⁷⁹ Then, on March 8, 2021, Practice Fusion's former Director of National Accounts, Steven Mack, pled guilty to one count of attempting to obstruct a federal investigation into the relationship between Practice Fusion and Purdue.⁵⁸⁰

vii. Another Mea Culpa

764. "I was horrified."

765. So wrote Ryan Howard, co-founder and former Chairman and CEO of Practice Fusion, describing his reaction to "encountering some staggering news:" "Practice Fusion, a company I founded in 2005 and left in 2015, had reached a settlement with the DOJ for suddenly partnering with an opioid manufacturer, and for encouraging doctors on the platform to prescribe opioids to patients." ⁵⁸¹

766. Howard "founded [Practice Fusion] in 2005 and was its Chairman and CEO until 2015... Under Ryan's leadership the company raised \$134 million in capital Kleiner Perkins Caulfied & Byers... Deerfield Management Company [and others] to fuel its rapid growth." ⁵⁸²

767. Howard noted that Practice Fusion "advertised to doctors 230 million times to prescribe opioids," and concluded that his "successors" fell "prey to the allure of capital over human lives." He minced no words regarding the appropriate outcome for his "successors" at Practice Fusion for allowing the company to partner with Purdue: "Following my departure in 2015, my

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⁵⁷⁹ Press Release, "Justice Department Announces Global Resolution of Criminal and Civil Investigations with Opioid Manufacturer Purdue Pharma and Civil Settlement with Members of the Sackler Family," Department of Justice, October 21, 2020, *available at*: https://www.justice.gov/opa/pr/justice-department-announces-global-resolution-criminal-and-civil-investigations-opioid

⁵⁸² See https://www.practicefusion.com/practice-fusion-founders/

Fress Release, "Former Practice Fusion Sales Executive Pleads Guilty to Obstructing Government Investigations into Purdue Pharma and Practice Fusion," Department of Justice, March 8, 2021, available at: https://www.justice.gov/usao-vt/pr/former-practice-fusion-sales-executive-pleads-guilty-obstructing-government

⁵⁸¹ Ryan Howard, "How to Prevent Your Company from Being Used for Evil From a Founder Who's Been There," *Entrepreneur*, November 28, 2020, *available at*: https://www.entrepreneur.com/article/359624

⁵⁸³ Ryan Howard, "How to Prevent Your Company from Being Used for Evil From a Founder Who's Been There," *Entrepreneur*, November 28, 2020, *available at*: https://www.entrepreneur.com/article/359624 (emphasis in original).

successors seemingly adopted a new mission – they allowed the [Purdue] partnership to take root – and for this, I suggest they experience the full retribution of our justice system." ⁵⁸⁴

768. Howard suggested that other entrepreneurs and founders of companies look into registering as benefit corporations, or B corporations, in order to prevent their creation from "being used for evil." He also provided the following "full disclosure:"⁵⁸⁵

I was never contacted by the DOJ, or any other authority, regarding Practice Fusion's investigations. Likewise, I was not part of any conversation with any opioid manufacturer while I was CEO. The referenced partnership occurred in 2016, after my departure, as detailed in the official DOJ Report (p. 18).

769. Like the Sackler's before him, Howard desired to distance himself from Purdue Pharma. But, as noted above, Practice Fusion's relationship began in the fall of 2012, and Howard was CEO for approximately 2 years while the company partnered with and serviced Purdue. In the fall of 2012, Purdue was "working with Practice Fusion to conduct a pilot (test) with Butrans advertising on the EHR (electronic health record) site to determine if banner ads work at driving traffic to our e-marketing collateral (websites, savings cards online, patient education downloads, etc.). The strongest interest is in seeing if we can improve coupon and co-pay program utilization." 586

770.	And as early as September 13, 2012, Ryan Howard

586 PPLPC018000741102

⁵⁸⁴ Ryan Howard, "How to Prevent Your Company from Being Used for Evil From a Founder Who's Been There," *Entrepreneur*, November 28, 2020, *available at*: https://www.entrepreneur.com/article/359624

⁵⁸⁵ Ryan Howard, "How to Prevent Your Company from Being Used for Evil From a Founder Who's Been There," *Entrepreneur*, November 28, 2020, *available at*: https://www.entrepreneur.com/article/359624

77.1 Fil 25 2014					
771. Then, on February 25, 2014,					
772. In other words,					
Howard					
wrote, "I was not part of any conversation with any opioid manufacturer while I was CEO."589					
Maybe he forgot. ⁵⁹⁰ Howard's words of wisdom					
to other founders regarding preventing their companies being "used for evil" are heartfelt,					
nonetheless.					
773. On August 18, 2015, Ryan Howard stepped down from his role as CEO of Practice					
Fusion and became Chairman of the Board. The Chief Commercial Officer, Tom Langan, replaced					
Howard as CEO. Langan had joined Practice Fusion only one year before. ⁵⁹¹					
774. Within months of Howard's departure, Practice Fusion hired JPMorgan Chase to					
explore the possibility of the company's initial public offering at a valuation range of \$1.1 to \$1.5					
explore the possionity of the company is initial public offering at a variation range of \$1.1 to \$1.3					
587 PPLPC022000561019					
588 PPLPC022000361019 588 PPLPC021000631187 589 Ryan Howard, "How to Prevent Your Company from Being Used for Evil From a Founder Who's Been There,"					
Entrepreneur, November 28, 2020, available at: https://www.entrepreneur.com/article/359624					
590 Likewise, Howard may have forgotten See MNK-					
71_0006314898; MNK-T1_0006316021. 591 Mark Sullivan, "Practice Fusion CEO Ryan Howard steps down, becomes chairman of the board," <i>Venture Beat</i> ,					
August 18, 2015, <i>available at</i> : https://venturebeat.com/2015/08/18/practice-fusion-ceo-ryan-howard-steps-down-becomes-chairman-of-the-board/					

billion valuation in 2017. The valuation was based on projected annual revenues in 2018 of between \$155 million and \$181 million.⁵⁹² Its actual annual revenue in 2015, the year Practice Fusion engaged its financial advisors to explore sales options, was approximately \$50 million.⁵⁹³

775. Around the same time, however, Practice Fusion was seeking other options. Practice Fusion's board hired Evercore, an investment bank, in November 2015 to solicit interest from people who might want to buy Practice Fusion whole. Interested parties indicated a bid range for Practice Fusion between \$50 million to \$225 million, or around 0.5% of the IPO valuation.⁵⁹⁴

776. Four months after engaging Evercore and obtaining the low estimated offers, Practice Fusion was eager to show greater profitability as soon as possible so as to close the gap between the low valuations anticipated in a whole company sales transaction versus the billion dollar company it wanted to be in the event of an IPO.

777. In February 2016, Practice Fusion fired one quarter of its workforce. Howard's successor, Langan indicated the downsizing was necessary "to get the company to a profit, at the same time that the low-priced acquisition offers were starting to accumulate." ⁵⁹⁵

778. It was within this context, with Practice Fusion desperately seeking near term revenue in order to validate an inflated valuation of the company, that Practice Fusion partnered with Purdue, who was likewise willing to pay top dollar in order to obtain its *own* "near term" growth. The two companies needed each other. For example, in discussing the proposed CDS alert

⁵⁹² Katie Benner, "Practice Fusion Said to Hire JPMorgan Chase to Explore I.P.O.," *New York Times*, January 19, 2016, *available at*: https://www.nytimes.com/2016/01/20/business/dealbook/practice-fusion-said-to-hire-jpmorgan-chase-to-explore-ipo.html

⁵⁹³ Christina Farr, "Employees at Practice Fusion expected IPO riches, but got nothing as execs pocketed millions," *CNBC*, January 23, 2018, *available at*: https://www.cnbc.com/2018/01/23/practice-fusion-workers-got-nothing-in-deal-as-execs-made-millions.html

⁵⁹⁴ Christina Farr, "Employees at Practice Fusion expected IPO riches, but got nothing as execs pocketed millions," *CNBC*, January 23, 2018, *available at*: https://www.cnbc.com/2018/01/23/practice-fusion-workers-got-nothing-in-deal-as-execs-made-millions.html

⁵⁹⁵ Christina Farr, "Employees at Practice Fusion expected IPO riches, but got nothing as execs pocketed millions," *CNBC*, January 23, 2018, *available at*: https://www.cnbc.com/2018/01/23/practice-fusion-workers-got-nothing-in-deal-as-execs-made-millions.html

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⁵⁹⁶ Practice Fusion Information at Para. 45. ⁵⁹⁷ Christina Farr, "Practice Fusion, once headed for \$1.5 billion valuation, ends in 'disappointing' fire sale," CNBC,

million-in-cash.html

available at https://www.youtube.com/watch?v=YZ3ZjARBnrI

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⁶⁰⁰ Arthur Turner, Consulting is More Than Giving Advice, Harvard Business Review, September 1982, available at: https://hbr.org/1982/09/consulting-is-more-than-giving-advice.

January 8, 2018, available at: https://www.cnbc.com/2018/01/08/practice-fusion-acquired-by-allscripts-for-100-

⁵⁹⁸ See "Business Consulting at ZS: learn how ZS recruits and interviews talent," ZS Associates, March 16, 2018,

program with Purdue, Practice Fusion's Senior Vice President on May 11, 2015, noted that there was "urgency" for Practice Fusion to generate revenue. 596

Ultimately, Practice Fusion was sold to Allscripts, another EHR vendor, on January 779. 8, 2018, for \$100 million, more than 90% less than their projected IPO valuation, in "a disappointing fire sale."597

ZS – The Salesforce Specialists 0.

780. Management consulting is the business of providing solutions to corporate clients. "Business consulting is really focused on solving our clients' business problems, and it is a very diverse set of issues that we might tackle, for example, 'where are the growth the growth opportunities in our business, and how to we go after them,' to something very specific, like 'what is the comp design that we should put together to incentivize our sales representatives next quarter," explained Kelly Tousi, a Principal at ZS Associates. 598 "That's a wide range, as you can imagine, and we do everything in between," she said.⁵⁹⁹

781. Solutions take many forms, depending on and tailored to the client's needs. "Management consulting includes a broad range of activities, and the many firms and their members often define these practices quite differently."600

782. As described above, broadly speaking, there are two schools of management consulting, namely "Strategy Consulting" and "Implementation Consulting." ZS engages in both.

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783. ZS describes itself as "a professional services firm that works side by side with companies to help develop and deliver products that drive customer value and company results." "Impact where it matters," the ZS webpage declares. 602 ZS describes its impact as results, not just ideas. "That's why we partner with our clients from strategy to implementation and beyond." 603

784. Consistent with the origins of ZS in academia, the company's initial focus was on building models that could be used by their clients to drive decision-making regarding salesforce structure and operations. As the founders explain, "early in our modeling careers in the 1970's, our thinking was centered on models, and we believed that the model was a large and prominent art of solving sales-resource-optimization problems."

785. Like many consulting firms, ZS performs both strategy and implementation work for its clients. But what sets ZS apart is that it has developed a particular niche in offering these services in the context of pharmaceutical sales and marketing. ZS specializes in the optimization of pharmaceutical sales forces in order to maximize sales and profit. In fact, ZS boasts that, "[i]n its first three years, ZS helped eight of the 10 largest pharmaceutical companies in the world align territories and resize their sales forces. By 2011, ZS worked with 40 of the 50 largest drug makers in healthcare and 17 of the 20 largest medical device makers." 605

786. In addition to developing overall sales and marketing strategies for specific drugs and drug portfolios, ZS regards implementation work as a core component of its overall product offerings to its clients. Indeed, implementation – and in some instances wholesale outsourcing of key business functions to ZS – are included as a component of practically every project that ZS takes on for a client.

⁶⁰¹ See https://twitter.com/ZSAssociates.

 $^{^{02}}$ Id.

⁶⁰³ See https://www.zs.com/about/our-impact.

⁶⁰⁴ See Prabhakant Sinha and Andris Zoltners, Sales Force Decision Models: Insights from 25 Years of Implementation, Interfaces 31:3, Part 2 of 2, Pg. S35-36, May-June 2001.

⁶⁰⁵ See https://www.zs.com/about/our-story.

787. In the broadest of generalities, then, ZS' business model, as a provider of strategy and implementation consulting services to the pharmaceutical industry, is to partner with clients to pursue business objectives identified by ZS. Once the objective is identified, the client and ZS then engage in concerted action, as a seamless and cohesive unit, in order to implement the necessary means to achieve the objectives for the client.

788. Beyond these traditional consulting services, ZS Associates are also known to serve as thought leaders by authoring articles or providing soundbites or quotes in order to influence perceptions in their client's favor. For example, when physician's offices began restricting access to sales representatives, ZS joined with manufacturers to create content designed to counter this movement.

789. ZS optimizes pharmaceutical sales forces for the explicit purpose of increasing sales and profit for the manufacturer client. In 2001, the founders of ZS published a paper entitled "Sales-Force Decision Models: Insights from 25 Years of Implementation." Describing ZS specialist expertise, the founders stated, "Over 25 years, we have developed many sales-force and modeling insights through over 2,000 projects with several hundred selling organizations in over 50 countries ... Two to three percent of all of the field salespeople in the US have been touched by the results. The firms had pressing issues that required quick attention. Companies sought help when merging separate selling organizations, when launching new products, when facing deregulation, or when faltering in performance."

790. In 2013, Chris Wright, ZS' current Chief Executive Officer, explained to the *New York Times*: "There's a group of geeks, if you will, who are running the numbers and helping the

⁶⁰⁶ See George Chressanthis et. al., Can Access Limits on Sales Representatives to Physicians Affect Clinical Prescription Decisions? A Study of Recent Events With Diabetes and Lipid Drugs, The Journal of Clinical Hypertension, Vol. 14, No. 7, July 2012, available at: https://onlinelibrary.wiley.com/doi/epdf/10.1111/j.1751-7176.2012.00651.x

⁶⁰⁷ *Id.* at S9.

sales guys be much more efficient."⁶⁰⁸ The effect is "what would happen if Arthur Miller's Willy Loman met up with the data whizzes of Michael Lewis's 'Moneyball."⁶⁰⁹

791. Wright was only a managing director at ZS when he provided his comments to the *Times* in 2013. In his 25 years at ZS, according to an "Impact Fact" described on ZS' website, "Chris has helped dozens of pharmaceutical companies differentially resources their sales deployments, leading to multibillion-dollar industry cost savings."⁶¹⁰

792. By exploiting "vast databases of patient and doctor information," companies like ZS can provide advanced analytics capabilities to clients to maximize sales efforts. "They know whether patients are filling their prescriptions – and refilling them on time. They know details of patients' medical conditions and lab tests, and sometime even their age, income and ethnic backgrounds." 611

793. ZS cannot drive customer value or company results, however, if its work is placed in a drawer and ignored. As such, ZS does not merely provide advice or models to clients. Instead, it works "side by side" with them to achieve results the client alone cannot. This reality was recognized early on in the history of ZS. "Over the years, we have realized that we spend much more energy on other activities, such as articulating the issues, building databases, and dealing with change management and implementation. For example, in the geographic deployment work we have done, we spend over 95 percent of the time in activities *unrelated to model building*." 612

794. Likewise, clients of ZS do not wish to pay top dollar for specialized management consulting services regarding their crucial sales and marketing practices merely for suggestions

⁶⁰⁸ Katie Thomas, "Pills Tracked from Doctor to Patient to Aid Drug Marketing," *New York Times*, May 16, 2013, available at: https://www.nytimes.com/2013/05/17/business/a-data-trove-now-guides-drug-company-pitches.html ⁶⁰⁹ Id.

⁶¹⁰ See https://www.zs.com/about/our-people/Chris-Wright

⁶¹¹ Katie Thomas, "Pills Tracked from Doctor to Patient to Aid Drug Marketing," *New York Times*, May 16, 2013, available at: https://www.nytimes.com/2013/05/17/business/a-data-trove-now-guides-drug-company-pitches.html
⁶¹² *Id.* at S36.

about how best to do things on their own. Just as an automobile manufacturer does not hire an airbag company to advise it on how to design, create, and install safe airbags in its cars, but instead just hires the airbag company to sell it safe airbags; pharmaceutical manufacturers hire consultants like ZS to implement mission-critical salesforce tasks they do not have the capability to perform on their own.

ZS speaks in terms of "optimizing" its clients' efforts to sell its products. ZS' clients 795. are for-profit companies, and "optimization" implies a specific variable you are optimizing. In the case of ZS, the variable is the amount of money the client can make.⁶¹³

796. ZS applied its hard-won expertise to multiple clients, "optimizing" their salesforces for the purposes of maximizing the profits derived from selling controlled substances.

i. The Salesforce – Pharma's Engine

797. Sales forces are a major component of pharmaceutical companies' operations. Indeed, they are the core of the industry. In 2007, it was estimated that there were 100,000 pharmaceutical sales representatives in the United States pursuing approximately 200,000 prescribers.614

798. These armies of sales reps are employed by pharmaceutical companies and detailed to health care providers to market the companies' drugs to those with the power to prescribe them. By 2000, at the outset of the opioid crisis, pharmaceutical companies were spending in excess of \$15 billion annually promoting drugs, with 84% of the total spend directed at detailing sales representatives to prescribers, drug samples, and ads in medical journals. 615

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See https://www.zs.com/solutions/artificial-intelligence-and-analytics/analytics (Touting ZS's "Analytics 27 optimization" solutions, ZS emphasized, "You need to ensure that you're investing in the right analytics capabilities, tapping into the right data sets and maximizing your ROI with key insights that drive business transformations.")

⁶¹⁴ Tobias L. Milrood, When Drug Representatives Go Too Far, American Association for Justice, February 2007. 615 M.B. Rosenthal et al., Promotion of Prescription Drugs to Consumers, 346 New England J. Med. 498 (2002).

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799. "Because of the large size of pharmaceutical sales forces, the organization, management, and measurement of effectiveness of the sales force are significant business challenges." This is ZS' niche. ZS tells its clients how to optimize, incentivize, and deploy these armies of pharmaceutical sales representatives for the purpose of maximizing revenue. In the words of ZS' founders, "[t]hat marketing investment drives sales is a fundamental principle supported by data."617 ZS has observed the statistically significant relationship between sales force effort and sales of pharmaceuticals, as depicted in the following scatter plot⁶¹⁸:

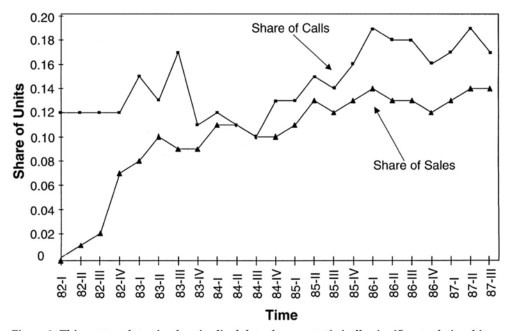


Figure 2: This scatter plot using longitudinal data shows a statistically significant relationship between sales-force effort and sales for a product sold by a pharmaceutical sales force. Every dot represents a quarter of the year.

800. ZS provided these services to numerous opioid manufacturers, which produced controlled substances known to be addictive at the time ZS advised them, for the explicit purpose of maximizing the sales and revenue of these deadly and addictive drugs during the pendency of a nationwide opioid crisis wrought by the over-selling of opioids by ZS' clients.

⁶¹⁶ Tobias L. Milrood, When Drug Sales Representatives Go Too Far, American Association for Justice, February

⁶¹⁷ Prabhakant Sinha and Andris Zoltners, Sales Force Decision Models: Insights from 25 Years of Implementation, Interfaces 31:3, Part 2 of 2, Pg. S10, May-June 2001. Id. at S10. ⁶¹⁸ *Id.* at S12.

ii. ZS – Pharma's long-term partner

801. Like many participants in the pharmaceutical consulting space, ZS does not merely provide advice to its clients on a one-off basis. Rather, according to ZS, a client relationship is "about results, not just ideas. That's why we partner with our clients from strategy to implementation and beyond." "We work side-by-side with you at every stage to help you achieve success." These partnerships are *long term*. For instance, ZS' founders have observed that, "having worked with some managers *repeatedly for over a decade or more*, we have observed patterns in the ways managers use consulting assistance and models." ⁶²¹

802. At ZS, "business consulting is really focused on solving our clients' business problems, and it is a very diverse set of issues that we might tackle, for example you could be anything from 'where are the growth opportunities in our business and how should we go after them,' to something very specific, like 'what is the comp design that we should put together to incentivize our sales representatives next quarter'? So that's a wide range, as you can imagine; we do everything in between."⁶²²

803. In other words, ZS does not merely provide ideas and advice to its clients. Rather, it designs and implements sales force models aimed at the most efficient allocation of marketing spending and maximization of profits for their manufacturer clients. In fact, ZS' recommendations are routinely implemented on such a systematic and industry-wide basis that ZS itself is now able

⁶¹⁹ See https://www.zs.com/about/our-impact

⁶²⁰ *Id*.

⁶²¹ Prabhakant Sinha and Andris Zoltners, *Sales Force Decision Models: Insights from 25 Years of Implementation*, Interfaces 31:3, Part 2 of 2, Pg. S37, May-June 2001 (emphasis added).

⁶²² See ZS Associates, "Business consulting at ZS: learn how ZS recruits and interviews talent," available at: https://www.youtube.com/watch?v=YZ3ZjARBnrI. Incidentally, the factors ZS articulates in its sales pitch as firm competencies – identifying growth opportunities and incentivizing the sales representatives – are both identified as principal origin points of the opioid epidemic by Dr. Van Zee as early as 2002.

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to draw on its own history of implementing change for its clients, as a data set of its own worthy of academic study.⁶²³

iii. ZS services its clients

- 804. Even though the marketing of OxyContin has been described as the "taproot" of the opioid epidemic, Purdue was not the only manufacturer to zealously market their own opioids. Nor was Purdue ZS' only opioid client.
- 805. ZS provided similar services, as explained below, to fellow opioid manufacturers Mallinckrodt Pharmaceuticals, Endo Pharmaceuticals, Teva Pharmaceuticals, and Johnson & Johnson's Janssen Pharmaceuticals. Consistent with its work for Purdue, ZS designed, implemented, and optimized salesforce strategies to maximize the profits derived from selling a controlled substance for practically every major opioid manufacturer. ZS' client work is described in individual detail below.

1. Purdue

806. As described above⁶²⁴, after 2007 Purdue was pursuing the Sackler's goal of maximizing near-term OxyContin sales so as to extract as much money from Purdue as possible in order to diversify away from the dangerous "concentration of risk" that continued financial reliance on Purdue represented for the billionaire family.

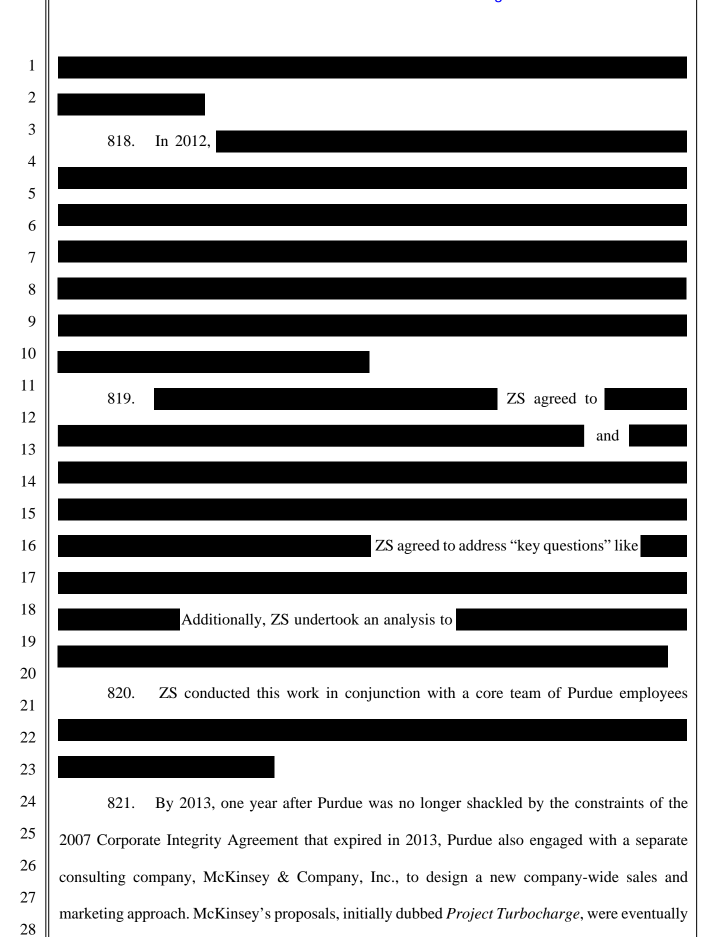
807. ZS happily agreed to help, and by ZS and Purdue were working together to increase sales of Purdue's opioids.

based on ZS' own independent research and unique methodologies, including modelling expertise. Purdue adopted ZS' strategies and worked closely with ZS to

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⁶²³ Prabhakant Sinha and Andris Zoltners, *Sales Force Decision Models: Insights from 25 Years of Implementation*, Interfaces 31:3, Part 2 of 2, Pg. S18-19, May-June 2001 ("The repeated application of several normative sales-forcedecision models has produced a series of insights that have led to a number of valuable sales-force insights.") ⁶²⁴ *See* Paragraphs 87-93, *supra*.

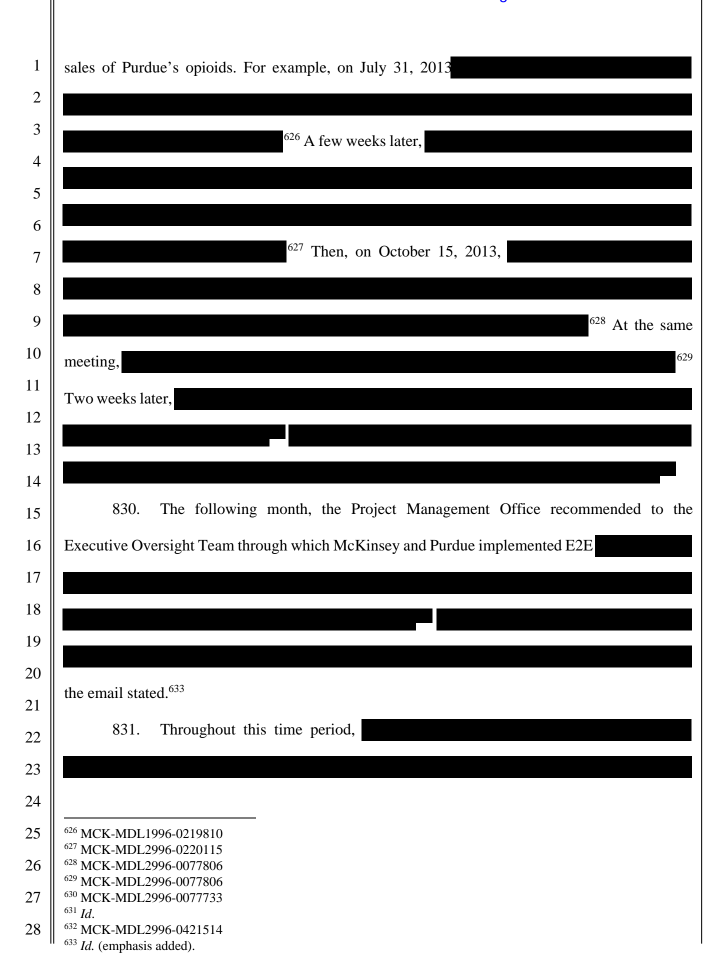
1	Workshops" with Purdue's staff regarding the best ways to undertake the changes in strategy
2	identified by ZS.
3	813. In
4	while Purdue was still bound by the Corporate Integrity
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6	Agreement. This time ZS' work related to
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8	814. The work took a holistic approach to Purdue's entire sales and marketing efforts for
9	its pain portfolio, including OxyContin, MS Contin, and RyzoltTM.
10	815. In addition to working on Purdue's existing pain portfolio, in ZS assisted
11	Purdue in
12	At that early point in the opioid crisis, Purdue was already
13	
14	interested in expanding into products for the treatment for opioid use disorder, which, according to
15	ZS,
16	816. ZS focused on answering mission-critical questions for Purdue, including but not
17	limited to:
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24	817. As always, ZS' work included an implementation component, including
25	and
26	ZS assured Purdue that
27	Zis assured Furdue that
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rechristened *Evolve to Excellence* and were implemented by McKinsey and Purdue for the explicit purpose of maximizing opioid sales despite the by-then obvious risks associated with selling as much OxyContin as possible.

- 822. ZS worked in cooperation with McKinsey and Purdue to implement and continually refine *Project Turbocharge*, including McKinsey's efforts to target the highest prescribers of OxyContin and blitz them with the newly turbocharged sales force. ZS worked with an Executive Oversight Team and Project Management Office, comprised of Purdue and McKinsey staff, to implement McKinsey's plans for Purdue.
- 823. That same year, despite significant headwinds, OxyContin sales finally peaked. The restrictions on Purdue's sales and marketing methods contained in the Corporate Integrity Agreement should have resulted in fewer overall OxyContin sales. Within five years of Purdue's guilty plea, however, OxyContin sales tripled.
- 824. ZS played a crucial role in accomplishing this feat. It presented specific plans to Purdue, which Purdue adopted and spent hundreds of millions of dollars implementing alongside ZS and other consultants. The result: a final spasm of OxyContin sales before the inevitable decline of the drug.⁶²⁵
- 825. In August of 2013, McKinsey urged, as part of the overall sales maximization approach, that sales representatives should devote two-thirds of their time to selling OxyContin and one-third of their time selling Butrans, another Purdue product. Previously, the split had been fifty-fifty.
- 826. Two months later, ZS sought to answer the question,

⁶²⁵ On February 10. 2018, Purdue announced that it is no longer marketing opioids, and disbanded its OxyContin sales force.



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634 MCK-MDL2996-0326142 (emphasis added).

By this time, ZS had also begun providing its AccessMonitor software for Purdue 832. to use. AccessMonitor is one example of a common tactic used by consultants to maintain an ongoing revenue stream from its clients, separate and apart from traditional project-based work: the development and marketing of "leave-behind" products, such as software applications, that are sold to clients as tools that can be used by the business on an on-going and recurring basis, separate and apart from the project-based consulting work that is ZS' core offering.

833. As described by famed Harvard Business School Professor Clayton Christensen, these sorts of "software and technology-based analytics and tools that can be embedded at a client," are a tool used by a consultancy to deepen its partnerships with clients and earn additional and recurring revenue from them. Tools such as ZS' AccessMonitor, Prof. Christensen noted, provide "ongoing engagement outside the traditional project-based model" traditionally used by consultants.635

By 2017, McKinsey was working on Project Scottsdale for Purdue. The goal of the project was to "transform Purdue's entire business model" by splitting Purdue's assets into three separate companies and downsizing the employee headcount at the companies by 500 people. 636 Just like Project Turbocharge, ZS collaborated with McKinsey on Project Scottsdale. In a December 2017 internal McKinsey email, McKinsey consultant Albert Lee reported to John Goldie

⁶³⁵ Clayton Christensen, Dina Wang, and Derek van Bever, "Consulting on the Cusp of Disruption," Harvard Business Review, October 2013, available at https://hbr.org/2013/10/consulting-on-the-cusp-of-disruption

⁶³⁶ The Firm and the FDA: McKinsey & Company's Conflicts of Interest at the Heart of the Opioid Epidemic, Interim Majority Staff Report, Committee on Oversight and Reform, U.S. House of Representatives, April 13, 2022, at Pg. 27, available https://oversight.house.gov/sites/democrats.oversight.house.gov/files/2022-04-13.McKinsey%20Opioid%20Conflicts%20Majority%20Staff%20Report%20FINAL.pdf

and Amir Golan that work on re-sizing Purdue's sales forces pursuant to the goals of Project Scottsdale was "a WIP [work in progress] with ZS."637

- 835. In December of 2015, ZS' client Purdue agreed to a settlement with the State of Kentucky relating to the improper marketing of OxyContin and other Purdue products. Purdue agreed to pay \$24 million in conjunction with the settlement.
- 836. Despite this second enforcement action against its client, ZS' work on Purdue's sales and marketing efforts continued unabated. Throughout its relationship with Purdue, ZS worked on core functions of Purdue's efforts to sell its drugs.
- 837. These core functions were previously identified as particular areas of concern with respect to Purdue's business conduct, and were specifically monitored and regulated under the 2007 Corporate Integrity Agreement, which governed, *inter alia*:
 - "selling, marketing, promoting, advertising, and disseminating Materials
 or information about Purdue's products in compliance with all applicable
 FDA requirements, including requirements relating to the dissemination
 of information that is fair and accurate ... including, but not limited to
 information concerning the withdrawal, drug tolerance, drug addiction
 or drug abuse of Purdue's products;
 - compensation (including salaries and bonuses) for Relevant Covered Persons engaged in promoting and selling Purdue's products that are designed to ensure that financial incentives do not inappropriately motivate such individuals to engage in the improper promotion or sales of Purdue's products; ...
 - the process by which and standards according to which Purdue sales representatives provide Materials or respond to requests from HCP's [health care providers] for information about Purdue's products, including information concerning withdrawal, drug tolerance, drug addiction, or drug abuse of Purdue's products," including "the form and content of Materials disseminated by sales representatives," and "the internal review process for the Materials and information disseminated by sales representatives." 638

⁶³⁷ MCK-MDL2996-0334687

⁶³⁸ See https://s3.documentcloud.org/documents/6452110/2007-Purdue-Corporate-Integrity-Agreement.pdf at pgs. 7-9.

838. In fact, under the terms of Paragraph II.C.1(b) of the Corporate Integrity Agreement, ZS, as a contractor to Purdue performing sales and marketing functions for the company, was a "Covered Person" subject to the strictures of the CIA.⁶³⁹

- 839. In addition to ZS' "expertise and thought leadership," ZS' assumption of these obligations for Purdue involved the deployment of Javelin, its proprietary salesforce optimization software tool. Like AccessMonitor, described *supra*, Javelin was a software tool that ZS could embed with clients. With Javelin, a ZS client can "streamline sales performance management with a comprehensive platform that simplifies sales strategy management and helps you build and motivate a successful sales force." The Javelin products include "suites" of software for the management and operation of Incentive Compensation (IC) and Call Planning (CP) functions. 641
- 840. Upon information and belief, all of Purdue's then 712 sales representatives were licensed users of ZS' trademarked Javelin suite of software solutions.
- 841. On October 20, 2020, Purdue entered into a plea agreement with the United States Department of Justice to plead guilty to improper marketing of OxyContin and other opioids, again.⁶⁴² This time the plea agreement concerned conduct from 2010 to 2018. ZS collaborated with Purdue on its sales and marketing practices the time period relevant to Purdue's second guilty plea.⁶⁴³

⁶³⁹ The relevant language in the CIA provides: "Covered Persons" includes ... all contractors, subcontractors, agents, and other persons who perform sales, marketing, promotional, pricing, government contract, or regulatory functions ... on behalf of Purdue." *Id.* at 2.

⁶⁴⁰ See https://www.zs.com/products/javelin.

⁶⁴¹ *Id*.

⁶⁴² See https://www.justice.gov/opa/press-release/file/1329576/download

on February 10, 2018, Purdue announced that it is no longer marketing opioids, and disbanded its OxyContin sales force. "OxyContin maker stops promoting opioids, cuts sales staff," *Reuters*, February 10, 2018, *available at*: https://www.reuters.com/article/us-usa-opioids-purduepharma/oxycontin-maker-stops-promoting-opioids-cuts-sales-staff-idUSKBN1FU0YL ("OxyContin maker Purdue Pharma LP said on Saturday that it has cut its sales force in half and will stop promoting opioids to physicians, following widespread criticism of the ways that drugmakers market addictive painkillers."). As such,

842. Purdue agreed to plead guilty to a dual-object conspiracy to defraud the United States and violating the Food, Drug, and Cosmetic Act, 21 U.S.C. § 331, 353, among other charges, relating to its opioid sales and marketing practices after the 2007 guilty plea.⁶⁴⁴ Purdue's coconspirators were not identified in the plea agreement.

843. Purdue's second guilty plea concerns Covered Conduct (as defined in the plea agreement) relating to Purdue's sales and marketing efforts that directly implicates ZS in the conspiracy. ZS' work for Purdue described in this Complaint was a core component of the sales and marketing tactics that lead to Purdue's second guilty plea.

2. Mallinckrodt

844. Upon information and belief, around the same time ZS was working with Purdue to implement McKinsey's *Project Turbocharge* to maximize OxyContin sales by continual refinement of physician targeting and other sales and marketing tactics, ZS was also working with another long-term client, Mallinckrodt,

845. Mallinckrodt is the largest supplier of opioid pain medications and among the top ten generic pharmaceutical manufacturers in the United States. In 2015, for instance, Mallinckrodt's opioids accounted for approximately one quarter (25%) of the entire annual production quota for controlled substances under DEA regulations. Mallinckrodt produced the following branded and generic opioids:

⁶⁴⁴ See https://www.justice.gov/opa/press-release/file/1329576/download

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Product Name	Chemical Name			
Exalgo	Hydromorphone hydrochloride, extended release			
Xartemis XR ¹⁰⁴	Oxycodone hydrochloride and acetaminophen (extended release)			
Roxicodone ¹⁰⁵	Oxycodone hydrochloride			
Generic	Oxymorphone hydrochloride (extended release) (generic Opana)			
Generic	Oxycodone (extended release) (generic OxyContin)			
Generic	Morphine sulfate, extended release			
Generic	Morphine sulfate oral solution			
Generic	Oxycodone and acetaminophen (Percocet)			
Generic	Hydrocodone bitartrate and acetaminophen (Vicodin)			
Generic	Hydromorphone hydrochloride			
Generic	Hydromorphone hydrochloride, extended release (Generic Exalgo)			
Generic	Fentanyl transdermal system			
Generic	Oral transmucosal fentanyl citrate			
Methadose	Methadone hydrochloride			
Generic	Methadone hydrochloride			
Generic	Buprenorphine and naloxone			

846. Previously,

Mallinckrodt sought to use ZS' expertise to

and to

- 847. Mallinckrodt promoted Exalgo as having characteristics that made the drug less likely to be addictive or abused, despite the lack of FDA approval for the drug as "abuse-deterrent."
- 848. Then, on March 12, 2014, Mallinckrodt obtained FDA approval for Xartemis XR, its extended-release opioid tablet.⁶⁴⁵ Upon information and belief, by the time Mallinckrodt obtained approval to market its opioid, ZS had already established a long-term working relationship with Mallinckrodt regarding the sales and marketing of Mallinckrodt's portfolio of pain medications.
 - 849. ZS was intimately involved from the outset.

⁶⁴⁵ See "Xartemis receives approval: May reduce opioid abuse," Formulary Watch, March 28, 2014, available at https://www.formularywatch.com/view/xartemis-xr-receives-fda-approval-may-reduce-opioid-abuse

850. In anticipation of Xartemis' launch, Mallinckrodt augmented its salesforce by adding hundreds of contracted sales representatives to promote the drug. CEO Mark Trudeau anticipated Xartemis would generate "hundreds of millions" in revenue for Mallinckrodt.⁶⁴⁶

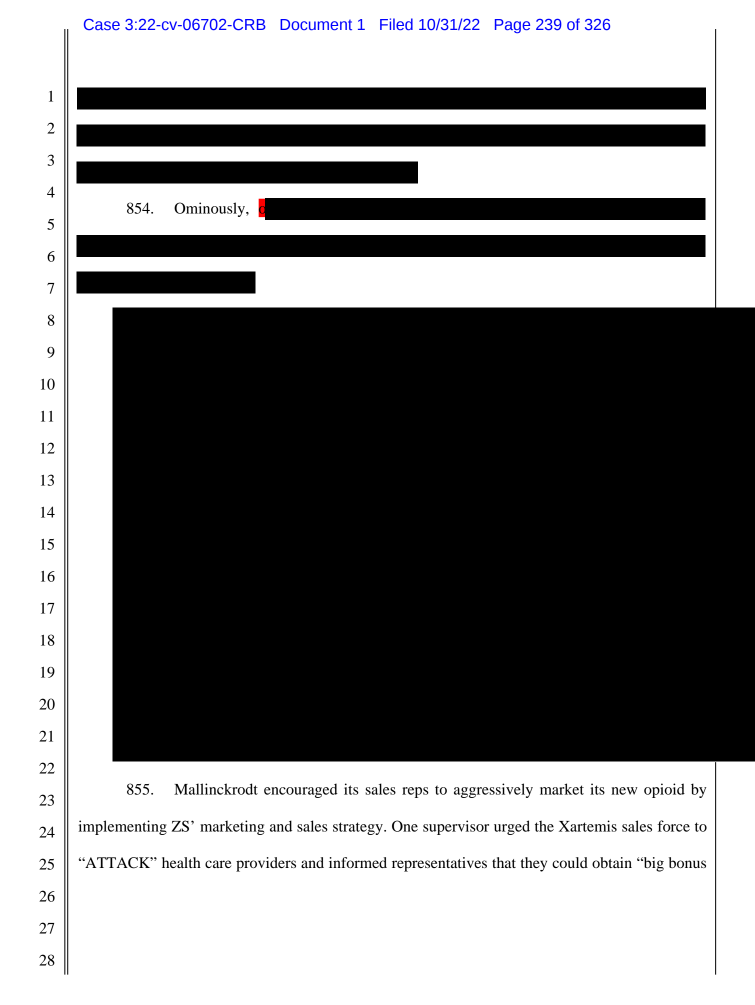
851. In September, only months after FDA approval, ZS unveiled an overall sales and marketing strategy for the drug. The scope of the plan was all-encompassing, including overall global strategy as well as granular details of execution and implementation. Tactics to be deployed in ZS' plan included the deployment of marketing materials in the offices of health care providers (including "in-office patient education materials" meant for the consuming public), physician targeting and decile segmentation, a video series interviewing clinicians about the benefits of the drug, patient testimonials, a speaker program, patient co-pay cards, targeting promotion outreach to regional associations of physician assistants⁶⁴⁷, and other efforts to maximize sales and revenue. Many of these same tactics were weapons in Purdue's arsenal, designed by ZS.



⁶⁴⁶ See https://www.bizjournals.com/stlouis/blog/health-care/2014/01/mallinckrodts-new-drug-should.html

⁶⁴⁷ This focus on physician's assistants is consistent with ZS' finding in an August 2016 report it issued, arguing that "[a]nother way to increase reps' chances for success is to expand the potential audience beyond the physician." ZS explained, "Other people on staff at the physician's office could be worthwhile targets for pharmaceutical messaging, such as nurse practitioners and physician assistants. *On average, people in these roles are incrementally more accessible than physicians*." *Physicians Becoming More Restrictive About Rep Sales Calls; Digital Communications Picks Up*, 28 No. 8 FDA Advertising & Promotion Manual News. 8, October 2016 (emphasis added).

⁶⁴⁸ "TRx" is an abbreviation for the measurement of increase in "total prescriptions," meaning all new prescriptions **plus** refills on those prescriptions. "Growing TRx," in other words, means encouraging refills of new prescriptions. This approach to maximizing revenue by encouraging refills of controlled substances known to be addictive carried obvious risks.



dollars" by waiting in front of doors of health care providers and using free trial offers to gain prescriptions. 649

- 856. Another sales supervisor encouraged sales representatives that Xartemis "is the BEST opportunity to make lots of money!!"⁶⁵⁰
 - 857. ZS designed and implemented the incentive compensation plan for Xartemis.
- 858. The above is even more alarming in light of the fact that ZS had been using its hublike position among many manufacturers to develop a business intelligence tool by analyzing call note reports from over 45,000 sales reps across the country. Given their ability to analyze the large amounts of data ZS possessed, ZS were the only ones with the keys to unlock the prescribing practices of each target physician.
- 859. Mallinckrodt repeatedly promoted Xartemis ZR as having physical properties that made the drug less likely to be addictive or abused, even though the drugs had never been approved by the FDA as abuse-deterrent. For instance, promotional materials provided to prescribers stated "Xartemis XR has technology that requires abuses to exert additional effort to extract the active ingredient from the large quantity of inactive and deterrent ingredients." 652
- 860. Not only had the FDA not approved Xartemis as an abuse-deterrent formulation, the marketing claims were false and misleading in that none of the characteristics of Xartemis address the most common form of opioid abuse: simple oral ingestion, swallowing the pill.

⁶⁴⁹ William K. Rashbaum, Roni Caryn Rabin and Danny Hakim, "Opioid Sales Reps Swarmed New York at Height of Crisis," *New York Times*, April 19, 2019, *available at* https://www.nytimes.com/2019/04/11/health/opioids-sacklers-

new-york-purdue.html

⁶⁵⁰ Id.

⁶⁵¹ See "Crossing the threshold: More than half of physicians restrict access to sales reps," *ZS Associates*, September 1, 2015, *available at*: https://www.zs.com/about/newsroom/crossing-the-threshold-more-than-half-of-physicians-restrict-access-to-sales-reps
28 652 *See* "Xartemis receives approval: May reduce opioid abuse," *Formulary Watch*, March 28, 2014, *available at*

⁶⁵² See "Xartemis receives approval: May reduce opioid abuse," Formulary Watch, March 28, 2014, available at https://www.formularywatch.com/view/xartemis-xr-receives-fda-approval-may-reduce-opioid-abuse

861. As stated above, the data and analytics capabilities of ZS, when tasked to the purpose of maximizing drug sales and revenue, should result in the unearthing of suspicious orders worthy of reporting to the DEA.⁶⁵³ Despite ZS working with Endo to assiduously study in granular detail the topic of where its pills are sold (and how to sell more of them), Endo nonetheless failed in its reporting obligations under the Controlled Substances Act.

862. In a July 2017 Memorandum of Agreement with the DEA and the Department of Justice, Mallinckrodt agreed that "at certain times during the Covered Time Period prior to January 1, 2012, certain aspects of Mallinckrodt's system to monitor and detect suspicious orders did not meet the standards outlined in letters from the DEA Deputy Administrator, Office of Diversion Control to registrants [Mallinckrodt] dated September 27, 2006, and December 27, 2007."654

3. Endo

863. ZS' relationship with Endo Pharmaceuticals

864. Upon information and belief,

months after Endo's launch of Opana ER, its extended-release oxymorphone tablet in 2006. At the time, Endo's marketing efforts were primarily focused on Opana ER, its branded extended-release oxymorphone hydrochloride tablet. Oxymorphone hydrochloride is three times as strong as morphine.

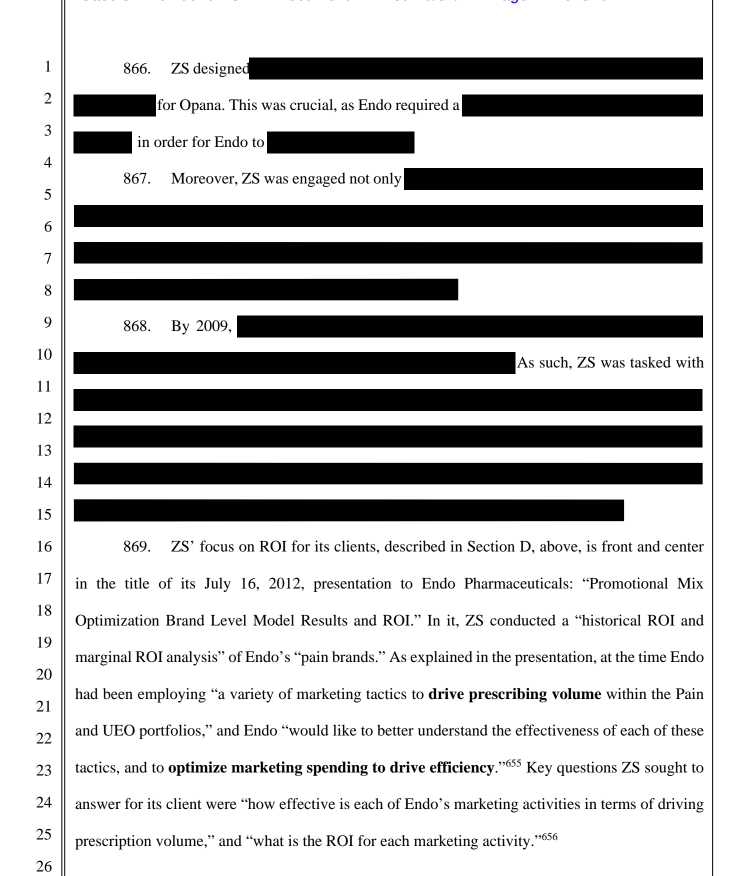
865. ZS' first known work for Endo involved developing and implementing

ZS

acknowledged.

 $^{^{653}}$ See supra, \P 64-65.

⁶⁵⁴ See the July 2012 Memorandum of Agreement between Mallinckrodt and the DEA at https://www.justice.gov/usao-edmi/press-release/file/986026/download



⁶⁵⁵ ZS Presentation to Endo dated July 16, 2012, In re: National Prescription Opiate Litigation, Case No. 1:17-md-02804-DAP, Doc. 2421-2, filed August 15, 2019 (N.D. OH).
656 Id.

⁶⁵⁷ *Id*.

⁶⁵⁸ *Id*. ⁶⁵⁹ *Id*.

870. ZS informed Endo plainly that, regarding its portfolio of pain medications, "[s]ales force detailing is the most impactful tactic, detailing accounts for 35-65 & of all sales and marketing impact."657 With respect to Opana, Endo's extended-release opioid tablet meant to compete with Purdue's OxyContin, sales force detailing accounted for 11.7% of the contribution to annual sales and profitability among Endo's sales tactics for Opana. In second place was co-pay cards, with 4.4%:658

% Contribution to annual sales and profitability by promotional tactic for each brand

Modeling time frame	Apr'11- Mar'12	Jan'11- Dec'11		Apr'11- Mar'12	
Tactic	LIDODERM	OPANA	OPANA Voltaren Gel		
SF Detailing	8.3%	11.7%	5.9%	8.2%	
NPP	3.1%	0.0%*	3.5%	7.5%	
Samples	5.0%	N/A	2.1%	1.0%	
Website	2.0%	2.1%	0.0%*	0.1%	
Journals	0.5%	0.3%	N/A	N/A	
Copay cards	0.4%	4.4%	2.7%	6.4%	
Speaker programs	Not planned for 2012	0.2%	N/A	N/A	
ALL TACTICS	19.3%	18.7%	13.2%	23.2%	
CARRYOVER	77.1%	73.4%	70.8%	45.2%	
OTHER FACTORS	3.6%	7.9%	15.0%	31.6%	
Positive mROI	Breakeven mROI	* Not a statistically si	100000	31.373	
ZS Associates		-9-	2013	07 16 Promo Mia Optimization - ROJ	

871. Opana ER constituted \$383 million in annual sales for Endo at the time of the ZS Associates analysis, good enough for the second highest selling Endo product analyzed by ZS. Of that \$383 million in sales, ZS determined that 19%, or \$72 million, is "driven by Sales and Marketing."659 Furthermore, ZS informed Endo that "most of the promotional channels for Opana

ER have high ROI." These "promotional channels" include marketing tactics such as copay cards, website advertising, medical journal sponsorship, speaker programs, and detailing to prescribers.

872. The Opana marketing messages whose delivery ZS sought to optimize for Endo conveyed misrepresentations regarding the dangers of the drug. For example, Endo maintained until April 2012 the website opana.com, which stated, "[m]ost healthcare providers who treat patients with pain agree that patients treated with prolonged opioid medicines usually do not become addicted." Upon information and belief, Endo did not conduct and does not possess data or evidence to support that statement. Furthermore, the statement is misleading in that it suggests that addiction is not a risk from taking extended-release opioids such as Opana.

873. Thus, ZS worked with Endo for at least a decade optimizing and implementing misleading sales and marketing tactics, *including managing critical sales and marketing functions* such as IC and CP. ZS' successful performance of these tasks was critical to Endo's efforts to market Opana. Without ZS, Endo could not have achieved its opioid revenue goals.

874. On March 3, 2016, the New York Attorney General announced a settlement with Endo to address the misleading marketing of Opana. The settlement required Endo "to cease all misrepresentations regarding properties of Opana ER, to describe accurately the risk of addiction to Opana ER, and to summarize studies regarding Opana ER on its website." The settlement further required Endo to create programs to prevent its sales representatives "from promoting [Opana ER] to health care providers who may be involved in the abuse and illegal diversion of opioids. Those safeguards were not in place under the sales and marketing program ZS designed and helped Endo implement for years. Even at the time of the settlement announcement, ZS was still working with Endo to shape and manage its Opana ER sales force.

⁶⁶⁰ See https://ag.ny.gov/press-release/2016/ag-schneiderman-announces-settlement-endo-health-solutions-inc-endo

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In May 2017, an advisory committee to the Food and Drug Administration recommended that Opana ER be withdrawn from the market due in part to the fact Opana ER could be "readily prepared for injection" (thereby bypassing the purportedly "abuse-deterrent" features of the formulation that Endo touted in its marketing) and was associated with outbreaks of HIV and a blood-clotting disorder known as thrombotic thrombocytopenic purpura ("TTP"). On June 8, 2017, the FDA adopted the committee's recommendation.⁶⁶²

- 876. One month later, on July 6, 2017, Endo announced that it would agree to cease marketing and selling Opana ER altogether.
- 877. Just as was the case with Purdue, ZS was working with Endo on marketing its branded opioid product at the time that the company voluntarily ceased selling and marketing the drug in response to the dangers of continuing to market it.

Teva

878. Not to be left out, Teva Pharmaceuticals also relied on ZS

Fentora is a fentanyl buccal tablet that is, "used for the treatment of breakthrough pain in cancer patients receiving opioid treatment and who have become tolerant to it."664 It was approved by the Food and Drug Administration in 2006 for this limited use, but, as the FDA noted two years later, "off-label prescribing has, unfortunately, been widely practiced." 665

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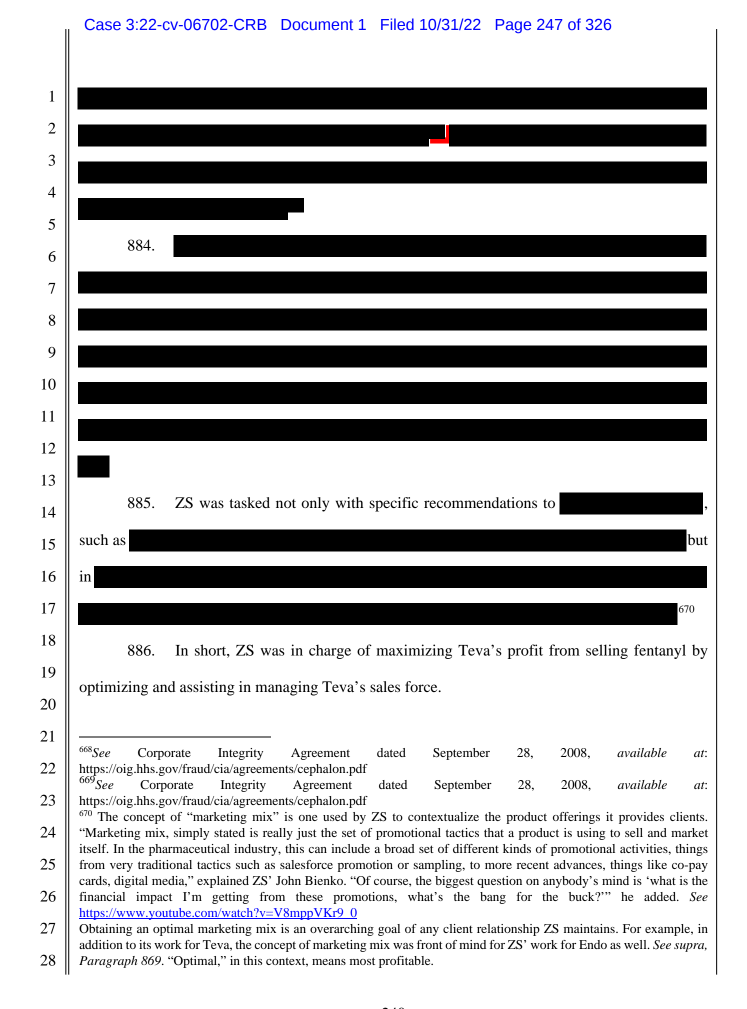
662 "FDA requests removal of Opana ER for risks related to abuse," June 8, 2017, available at: https://www.fda.gov/NewsEvents/Newsroom/PressAnnouncements/ucm562401.htm

⁶⁶³ See Teva Completes Acquisition of Cephalon, Fierce Pharma, October 11, 2011, available at: https://www.fiercepharma.com/pharma/teva-completes-acquisition-of-cephalon

Fentanyl Buccal Fentora) **Tablets** (marketed as Information, available at: https://www.fda.gov/drugs/postmarket-drug-safety-information-patients-and-providers/fentanyl-buccal-tabletsmarketed-fentora-information.

Memorandum from Bob A. Rappaport, MD to the Anesthetic and Life Support Drugs Advisory Committee (ALSDAC) dated April 26, 2008, Doc. 2231, In re National Prescription Opiate Litigation, No. 1:17-md-02804-DAP (N.D. Ohio) (filed August 13, 2019).

⁶⁶⁷ See https://www.fda.gov/drugs/information-drug-class/questions-and-answers-fda-approves-class-risk-evaluationand-mitigation-strategy-rems-transmucosal. - 239 -



5. Johnson & Johnson/Janssen

887. In addition to the manufacturer clients described above, ZS also worked for Janssen Pharmaceuticals, Johnson & Johnson's subsidiary that sold opioids

Johnson & Johnson used ZS just like everyone else: always.

888. The foregoing paragraphs make clear ZS' central role in the operations of its clients. Just as all pharmaceutical companies rely on IMS Health for sales data, or McKinsey for strategy consulting, practically *all* opioid manufacturers depend on ZS for salesforce optimization and sales and marketing advice. ZS is a critical part of the economic ecosystem that sells drugs in the United States.

p. Defendants work kills people.

889. For Defendants, the name of the game is pleasing clients. Client retention is a key metric by which publicly traded advertising firms like Publicis and Allscripts are judged in the capital markets and therefore pay particular attention to. Client retention ensures stable revenue streams for agencies like the Big Four, which is seen as more valuable in the eyes of capital markets than one-off engagements. At the same time, Defendants must exert effort (which costs money) to retain a new client. Client development is a *cost* for Defendants, and selling additional services to an existing client is an inexpensive way to maximize the return on the investment ("ROI") that Defendants make in developing and maintaining a client relationship.

i. ROI is King

890. Likewise, from the client's perspective, return on investment is the coin of the realm as well. The profit motive drives behavior on both sides. If an agency does not increase revenue and profit for the client, they are worthless.

891. The only thing that matters is the bottom dollar, and the bottom dollar is driven by the volume of goods sold.

892. Publicis understands this. In Publicis' own words:⁶⁷¹

In healthcare, performance cannot be measured in clicks or impressions or even office visits. Our only KPI is outcomes — business outcomes and patient outcomes. Our best-in-class outcomes analytics methodology enables us **to see real ROI**. But sound, evidence-based measurement isn't just about **proving** ROI — it's also about **optimizing** for it. With the best data-driven analytics team in healthcare media, we're able to achieve performance that outclasses the field — with **ROI four times greater than our closest competitor.**

- 893. Publicis' actions for its clients provide numerous real-world instances of this myopic focus on money. For example, in April 2013, Publicis implemented an email marketing campaign offering downloadable OxyContin "Savings Cards," which Publicis knew was a tactic for patient retention, or, keeping patients on OxyContin for longer periods of time. Again, Publicis couched the campaign's worth in terms of "return on investment" for Purdue. While Publicis charged Purdue \$47k for the campaign, Publicis and Purdue later assessed the incremental impact of the campaign to have been an additional 67,000 OxyContin prescriptions by April 2014.
- 894. Despite that apparent success, Publicis and Purdue wanted more. Notes from an internal Publicis meeting held four months later, in August of 2014, to discuss work on the Purdue account make it plain. Patient retention and return on investment remained top of mind:
 - Patient Retention "How do we get a 2:1 return on this?"
 - o What is a patient worth?
 - o What is a monthly script worth?
 - What is the value of getting just one more refill?
 - Isn't brand retention so much as overall portfolio retention
 - In the beginning, Purdue got reprimanded for going after patients too aggressively
 - Hence the focus is more company versus brand oriented; feels more altruistic
 - Determine at the patient level what they need for their condition
 - Does portfolio retention need to be veiled in a patient engagement program?

⁶⁷¹ See https://www.publicishealthmedia.com/our-approach/ (emphasis added).

"What is a patient worth?... How do we get a 2:1 return on this?... getting just *one* more refill." Internally, Publicis was direct in describing what matters.

895. The meeting was held to discuss additional projects to be performed in the realm of "patient retention" for Purdue. "Patient retention" for Purdue has all of the same benefits as "client retention" does for Publicis. In other words, Publicis was working to assure people stayed on their OxyContin prescriptions and never quit. Given the unpleasantness of that goal, Publicis wondered, "Does portfolio retention need to be veiled in a patient engagement program?"⁶⁷²

896. Likewise, ROI was paramount in Publicis' decision to engage Practice Fusion on the Purdue account. In a July 18, 2014, internal email, Publicis' John Dwyer asked a colleague, "Can you find out if they [Practice Fusion] have any kind of ROI of Rx impact metrics around these that they can share?"

897. ROI was king, but Publicis knew you couldn't just *say* that. In October 2014, Publicis was working with Purdue to devise an "Outward Facing Strategy," whereby Publicis would help Purdue present itself as a responsible corporate citizen to the world. Publicis noted that the current corporate tagline of "profitable growth" was not "outward facing" and was instead "strictly for internal use."

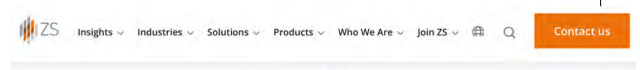
898. Describing the bottom line regarding the services ZS provides its clients, ZS Principal John Bienko stated, "Of course, the biggest question on anybody's mind is 'what is the financial impact I'm getting from these promotions, what's the bang for the buck?" "673

⁶⁷² For a discussion of why Publicis might wish to "veil" its retention goals in the guise of "patient engagement," *see supra.*, Section IV(e)(iii)(c).

⁶⁷³ See ZS Associates, "Marketing mix strategy: why it's so important for pharmaceutical marketing," available at: https://www.youtube.com/watch?v=V8mppVKr9 0

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899. ZS' raison d'etre is maximizing return on investment for all sales and marketing spending for a pharmaceutical manufacturer. ZS was not shy about couching its entire product line to pharmaceutical manufacturer clients in terms of ROI⁶⁷⁴:



WHAT WE DO

Challenges we solve



Identifying the best way to reach customers

To ensure that your message makes the desired impact, you need to select the right channels and messages based on observed customer preferences.



Creating realistic, actionable plans

Success is in the details. How can you create practical and actionable plans to deliver the right message in the right channel at the right moment to achieve optimal effect?



Striking the right message balance

What messaging are your taking to your target audience?

Does your content balance promotional priorities with

customer needs?



Assessing performance to improve ROI

To determine where your marketing investments will reap the best returns, you need to evaluate which promotional channels will have the greatest impact on brand performance.

- 900. ROI is the basis upon which ZS advertises and sells its services to pharmaceutical companies, and it is the principal reason that a pharmaceutical company hires ZS in the first place. As established herein, ZS couched substantially *all* of its proposals to work for opioid manufacturers in terms of how much money they will make by doing what ZS recommends.
- 901. This myopic focus on the bottom line, when applied to controlled substances known to be addictive, would have predictable consequences.
 - ii. McKinsey Publicis Knew; Practice Fusion Knew; ZS Knew; Everyone Knew

⁶⁷⁴ See https://www.zs.com/solutions/marketing/promotions-and-marketing-mix

902. As described above, the problem with broadly and aggressively marketing addictive opioids was apparent from the start: controlled substances are *controlled* precisely because they should not be sold to maximize volume and profits. Defendants actively disregarded this plain fact, repeatedly, and for decades.

1. McKinsey Knew

903. The deceptive marketing strategies McKinsey developed and helped to implement were successful. Its granular growth tactics, myopically focused on increased revenues for its clients, substantially contributed to an explosion in the use of opioids across the country. Approximately 20% of the population between the ages of 30 and 44, and nearly 30% of the population over 45, have used opioids. Opioids are the most common treatment for chronic pain, and as of 2016, 20% of office visits for non-cancer pain included the prescription of an opioid.⁶⁷⁵

904. The dangers of opioids were known to McKinsey at the time it engaged in the misconduct described in this Complaint. The addictive potential of opioids and the need for control and restraint in their use was internally understood, as was the likelihood of large-scale opioid addiction, abuse, overdoses, illness, and early death resulting from sharply increased use.

905. McKinsey also performed its own research in evaluating the anticipated effects of Project Turbocharge. An April 2014 implementation update observed an increase in sales calls, as well as that "OxyContin [health care providers] with increased calls consistently outperform HCPs with decreasing or no change in call frequency."

906. McKinsey continued working with Purdue long after the severity of the opioid crisis was well known. McKinsey knew that high dose OyxContin prescriptions carried a serious risk of overdose. In 2017, over half of Purdue's opioids prescriptions exceeded the ninety mg morphine

⁶⁷⁵ Deborah Dowell, Tamara M. Haegerich, and Roger Choi, *CDC Guideline for Prescribing Opioids for Chronic Pain – United States*, 2016, CDC (March 18, 2016), https://www.cdc.gov/mmwr/volumes/65/rr/rr6501e1.htm

equivalence threshold a day—the recommended maximal dose per the 2016 CDC Guideline for Prescribing Opioids for Chronic Pain.

- 907. Purdue's 2007 guilty plea put McKinsey on notice of Purdue's misconduct. By that time, McKinsey had access to public information indicating that OxyContin and other opioids pose significant risk of addiction and misuse.
- 908. McKinsey was well aware of the risks of OxyContin based on its extensive experience in the pharmaceutical industry, close collaboration with Purdue, and participation in the regulatory submissions for reformulated OxyContin.⁶⁷⁶
- 909. The first bullet point of Purdue's 2007 "Observations and Activities Requiring an [Abuse, Diversion, and Detection] Report" was "[a]n apparent pattern of an excessive number of patients for the practice type[.]"⁶⁷⁷ Thus, McKinsey knew or should have known that there was a higher risk of abuse and diversion among high-volume prescribers.
- 910. What is more, on September 13, 2013 McKinsey briefed Purdue on the ongoing concerns regarding OxyContin addiction and diversion among prescribers:

676 McKinsey

McK-MAAG-0118819 (email chain).

677 PPLPC010000033944.

Findings on messaging and positioning

PRELIMINARY

- Opioids overall are still viewed as effective and necessary class of painkillers, though side effects and addiction are concerns
- · Key themes from prescriber interviews on abuse deterrents include:
 - Prescriber awareness of abuse deterrence and label change is mixed
 - Opinions on impact/efficacy of abuse deterrence vary
- Most prescribers are concerned about abuse, but attempt to establish measures to protect themselves
- Concerns remain that technology does not address oral abuse
- Less Informed prescribers ask for additional information and education around abuse deterrent formulations
- Existing market research suggests that most physicians do not feel that
 reformulation positively impacts their prescribing behavior, and that diversion,
 abuse and regulatory concerns continue to weigh on prescribers

McKinsey & Company | 2

- 911. In a PowerPoint slide entitled "Findings on messaging and positioning," part of a presentation to Purdue entitled "OxyContin growth opportunities: Phase 1 Final Report: Diagnostic," McKinsey noted that "most prescribers are concerned about abuse," and that "most physicians do not feel that [OxyContin] reformulation positively impacts their prescribing behavior, and that diversion, abuse and regulatory concerns continue to weigh on prescribers."
- 912. In an August 2017 presentation, McKinsey recognized that the opioid epidemic was "triggered, in large part, by a massive increase in prescribed opioids in the early 2000's."
- 913. McKinsey's presentations to Purdue included extensive discussion of doctors' concerns about opioid misuse and side effects, demonstrating McKinsey's awareness of the dangers of opioids. Rather than working to limit these disastrous effects, McKinsey treated doctors' misgivings as obstacles to confront with new messaging.
- 914. Indeed, one reason that *Purdue* had knowledge that their own products were addictive and dangerous is because McKinsey told them.

If McKinsey was not aware of the adverse consequences of OxyContin, the drug it

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915.

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⁶⁸³ *Id.* at 357:22–358:6.

⁶⁸⁴ Gerald Posner, *Pharma*, pg. 524 (Simon & Schuster 2020).

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686 https://www.justice.gov/opa/press-release/file/1329571/download

921. Instead, McKinsey focused on increasing opioid sales for its clients, despite knowing the harmful consequences of doing so. In yet another indication that OxyContin sales should not be turbocharged: during McKinsey's work for Purdue, Purdue was unable to purchase product liability insurance to cover its practice of selling OxyContin.

922. The basic premise of McKinsey's work put it on notice of the harmful consequences that would ensure. It was tasked with advising a monoline manufacturer of opioids about sales and marketing practices for its addictive products while that manufacturer was bound by a five-year Corporate Integrity Agreement covering the very same opioid sales and marketing practices. In 2012, OxyContin accounted for 94% of Purdue's revenue. 684 As late as 2018, it remained 84% of Purdue's revenue. 685 According to the U.S. Department of Justice, "[f]rom 2010 to 2018, Purdue's profits were almost entirely driven by its success in selling OxyContin."686 In 2015 alone, it obtained \$3 billion in annual opioid sales—a four-fold increase from its 2006 sales of \$800 million.

923. McKinsey's mandate was to increase Purdue's opioid sales during a time when Purdue was obligated to restrict its previous marketing strategies because those strategies had caused the overprescribing of opioids and the inevitable consequences thereof. McKinsey's job was to counter the intended results of the Corporate Integrity Agreement; to devise strategies to sell as many pills as conceivably possible. Under McKinsey's tutelage, Purdue's growth continued its upward trajectory unabated, the Corporate Integrity Agreement notwithstanding.

Publicis Knew 2.

924. Like Publicis Touchpoint Solutions' work with Orexo, Publicis's Saatchi and Saatchi also worked the other end of the epidemic. In 2017, it sponsored New York Festivals Global Awards Young Globals competition, where the winning team would win an internship opportunity. The Young Globals is "the only college/portfolio competition for healthcare advertising." The competition called for "creative challenge briefs" to be submitted "for the (fictional) National Opioid Addiction Prevention Council and invites student entrants to develop a unique and compelling multi-channel experience (print, social media, digital, etc.) for their project Push Back on Opioid Abuse," in order to "raise awareness about opioid addiction."

- 925. The same year, the President of the United States declared the opioid crisis a National Health Emergency.
- 926. Of course, everyone was aware of the problem long before 2017. On June 11, 2014, Publicis' John Dwyer was informed that the DEA believed "an oversupply of painkillers is fueling the black market for both prescription opioids and heroin," and furthermore that 668 individuals in the State of Massachusetts alone died of opioid overdoses in a single year (2012). In an internal Publicis email, Dwyer conceded, "in the opioid market there are so many other factors to consider than just 'how can we increase sales of our product?""
- 927. Dwyer's expressed concern about "other factors to consider" when selling opioids is belied the by reality of Publicis actual work with Purdue, where "increasing sales of our product" was *literally* the goal Publicis were hired to pursue.⁶⁸⁹ When the rubber meets the road, ROI is the only metric that matters.
- 928. Dwyer himself was in a unique position of knowledge. As a Publicis colleague told him in July 2016, "I doubt there's a marketer that knows the opioid marketplace like you do. You

- 250 -

⁶⁸⁷See https://www.lbbonline.com/news/new-york-festivals-young-global-awards-open-for-entries

⁶⁸⁸ See https://www.lbbonline.com/news/new-york-festivals-young-global-awards-open-for-entries

⁶⁸⁹ See PPLPC018000873870

are speaking from a very secure place of knowledge in terms of RX targeting and who is driving RXs."

929. That knowledge was placed front and center in Publicis' pitch to the non-profit Partnership to End Addiction for work on the website drugfree.org. In the pitch, Dwyer and Razorfish Health Vice President Karl Tiedemann highlighted Publicis' deep knowledge of the opioid marketplace, and described its work for Purdue:

We've been immersed in the evolving national opioid medication dialogue going on between pharma companies, the government and FDA, the media, and the public via **inside access as a trusted and informed consulting partner**... We've been ingrained in the evolution of these issues for 6 years as other companies have come and gone; monitoring the progress made with HCP's and patients. (emphasis added).

930. The detailed scope of Publicis' involvement in the opioid industry that Tiedemann and Dwyer set forth in their pitch to *a nonprofit combatting opioid addition* is remarkable. On March 1, 2016, Dwyer jotted down fifteen reasons why Publicis was "the indisputably most knowledgeable and most experienced agency to help with drugfree.org:"

1 Karl Tiedemann[karl.tledemann@razorfishhealth.com]; Scott Reese[scott.reese@razorfishhealth.com] To: From: John Dwyer Sent: Tue 3/1/2016 12:02:17 AM (UTC-05:00) 2 Subject: RFH experience for Drugfree.org Some notes I jotted down summarizing our relevant experience in this area since 2010 that positions us as the indisputably most 3 knowledgeable and most experienced agency to help with drugfree.org. 4 Worked with Purdue Pharma since Feb 2010 2. Not just on the promotion of their opioid medications OxyContin, Butrans, Targiniq and Hysingla ER 5 3. Worked on unbranded pain education program and website Partners Against Pain; rebranded, reinvented entire site twice including content and resources for HCPs, patients and caregivers. 4. Worked on company branding and website PurduePharma.com in 2011 and again in 2015 6 5. Worked on Purdue's OxyContin REMS program materials for soft re-launch/re-brand of OxyContin for reformulation in Sept 2010 7 6. Agency selected to lead work creating branding and site content development for multi-pharma company-sponsored class wide ER and LA opioid REMS program in 2012 7. Developed strategy, messaging and brand campaign for OADP (opioids with abuse-deterrent properties) unbranded campaign 8 used on teamagainstopioidabuse.com; site features content created for 7 stakeholder groups: Physicians; Patients/caregivers; Pharmacists; Payors; Parents and communities; Police and law enforcement; Pharma companies 9 8. Collaborated for the above with various departments outside of product Marketing: Public Affairs; Patient Advocacy; Promotional Med Ed; and programs Safeguard My Meds; Rx Safety Matters; In the Face of Pain; RxPatrol 9. Contributed to promotion and dissemination of all of the above via multi-channel promotion including websites, email 10 campaigns, banner ads, convention materials, print ads, media buying, SEM (paid search), and SEO 10. Created the first oxycontin.com publicly accessible website including a Patient site with Patient content and materials 11 11. Participated in dozens of market research projects with hundreds of HCPs, payers, pharmacists and patients over 6 years 12. Attended pain and pain medicine conferences since 2010 12 13. Attended Advisory Boards with PCPs and multiple pain treating specialties 14. We've been immersed in the evolving national opioid medication dialogue going on between pharma companies, the government and FDA, the media, and the public via inside access as a trusted and informed consulting partner 13 15. We've been ingrained in the evolution of these issues for 6 years as other companies have come and gone; monitoring the progress made with HCPs and patients 14 15 John Dwyer SVP, Group Account Director 16 John.dwyer@razorfishhealth.com P 212.771.5428 M 917.797.9317 17 3. **Practice Fusion Knew** 18 931. In April 2014, an internal email between Practice Fusion employees counseled that 19 20 "indicating that Purdue influenced clinical decisions through sponsored money has legal 21 implications versus a marketing program where a banner can be displayed and influence a 22 prescribing behavior."690 23 Practice Fusion knew which side of the line their conduct fell on. In a March 23, 932. 24 2015, internal email, Practice Fusion discussed pitching CDS alerts to Purdue, and stated that 25 Purdue "has communicated that the average dosage of OxyContin is declining," because prescribers 26 27 28

⁶⁹⁰ Practice Fusion Information at Para. 27.

are "hesitant about using high dosages to combat pain for a variety of reasons, mostly, political pressure." Because of this "pressure," Purdue might use Practice Fusion to boost prescriptions... "Purdue is toying with the idea of using Pain Assessment tools with the provider at every visit and before every RX."

- 933. In September 2015, a Practice Fusion employee advised a Practice Fusion colleague in an internal email, "I understand that the Purdue proposal has shifted to a commercial focus and that marketing folks were in the room instead of outcomes... there are several things incorrect with this presentation/proposal from pricing to products. Please do not share. Just be aware..."⁶⁹¹ That same month, another Practice Fusion employee wrote in a separate internal email describing a meeting with Purdue: "[W]e were talking to product managers, and they could care less about RWE [real world evidence]. For them, this was all about marketing."⁶⁹²
- 934. Practice Fusion knew that Purdue was only concerned about increasing prescriptions. On May 11, 2016, a Practice Fusion employee observed that in a meeting with Purdue to discuss the development of the CDS alert, he kept "hearing the client [Purdue] revert back to 'Rx lift' as the primary objective of the program, this came up in the kickoff meeting and again during last week's meeting." Repeatedly, Purdue made it clear to Practice Fusion what mattered.
- 935. Practice Fusion knew that even *Purdue* had legal concerns about the CDS alerts program. Purdue asked that Practice Fusion provide an update on the results of the CDS Program, including whether "the CDS alerts change prescribing behavior" with respect to EROS. The meeting between Practice Fusion and Purdue occurred on December 14, 2016. A Purdue attorney at the meeting expressed reservations about the CDS program and noted that the program had not received appropriate legal review within Purdue.⁶⁹³

⁶⁹¹ Practice Fusion Information at Para. 58.

⁶⁹² Practice Fusion Information at Para. 60

⁶⁹³ Practice Fusion Information at Para. 110.

936. A post-hoc legal review of the CDS alert program was conducted by Purdue with Practice Fusion's input in December 2016 and January 2017. Despite the legal review, none of the conduct that led to Purdue's subsequent guilty plea and Practice Fusion's deferred prosecution agreement was altered or paused. The conduct continued until 2019.

4. ZS Knew

937. ZS was in a truly unique position, given its dominance of pharmaceutical sales and marketing consulting, practically all industry participants were its clients. While advising multiple industry participants regarding the sales of competing products (OxyContin and Opana, for instance) at the same time, ZS was in a position to know confidential information and trade secrets of these clients. Indeed, the contracts between ZS and its clients specify that ZS will have access to the clients' confidential and proprietary information. Given the nature of ZS' work, it cannot adequately perform its function for clients without that access.

938. ZS' clients were repeatedly subjected to enforcement actions for their work selling opioids both before and during the pendency of the ZS client relationship. For instance, in addition to its 2007 guilty plea with the United States Department of Justice ("DOJ"), Purdue Pharma settled with the State of Kentucky in 2015 for \$24 million.⁶⁹⁴ The settlement concerned similar conduct as the 2007 guilty plea, including the sales and marketing of Purdue's opioids. ZS was involved in this work.

939. Two years later, on July 11, 2017, another ZS client settled charges that it failed to report suspicious orders of opioids and for various recordkeeping violations. In this case, Mallinckrodt's failure to comply with DEA regulations regarding the sales of opioids resulted in a \$35 million payment to the DOJ.⁶⁹⁵

⁶⁹⁵See https://www.justice.gov/opa/pr/mallinckrodt-agrees-pay-record-35-million-settlement-failure-report-suspicious-orders.

⁶⁹⁴ *See* the December 22, 2015, settlement between the Commonwealth of Kentucky and Purdue Pharma Inc., *available* at https://ag.ky.gov/pdf news/purduepharmaoxycontin.pdf.

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940. The settlement agreement related to conduct between 2008 and 2011, during which time ZS was advising Mallinckrodt.

941. Because of these client relationships, ZS was in a unique position to know how the entire industry's opioid sales and marketing tactics were playing out, both in terms of return on investment for their individual clients, as well as overall market trends such as the rise of the opioid crisis. Endo may not have known the specifics of competitor Purdue's marketing efforts for OxyContin, just as Purdue may not have known the specifics of Endo's Opana plan. But ZS knew both, as well as what Teva and Mallinckrodt were doing with their own branded opioid sales and marketing efforts in real time.

5. **Everyone Knew**

942. The evidence of a direct link between increased opioids marketing and sales and increased opioid abuse was everywhere. A 2007 study found "a very strong correlation between therapeutic exposure to opioid analysesics, as measured by prescriptions filled, and their abuse." 696 McKinsey evidently understands this. In a September 2016 online article, McKinsey asserts that "[t]here is no doubt that more consistent use of best practices – across geographic areas, institutions, and clinicians - would provide tremendous help in combating the crisis" and describes certain examples of such practices as "successful in reducing prescribing." 697

943. There is a "parallel relationship between the availability of prescription opioid analgesics through legitimate pharmacy channels and the diversion and abuse of these drugs and

⁶⁹⁶ Theodore J Cicero et al., Relationship Between Therapeutic Use and Abuse of Opioid Analgesics in Rural, Suburban, and Urban Locations in the United States, 16.8 Pharmacoepidemiology and Drug Safety, 827-40 (2007), available at https://onlinelibrary.wiley.com/doi/10.1002/pds.1452.

⁶⁹⁷https://www.mckinsey.com/industries/healthcare-systems-and-services/our-insights/why-we-need-bolder-actionto-combat-the-opioid-epidemic

associated adverse outcomes." 698 The opioid epidemic is "directly related to the increasingly widespread misuse of powerful opioid pain medications."699

In a 2016 report, the CDC explained that "[o]pioid pain reliever prescribing has 944. quadrupled since 1999 and has increased in parallel with [opioid] overdoses." Patients receiving opioid prescriptions for chronic pain account for the majority of overdoses. For these reasons, the CDC concluded that efforts to rein in the prescribing of opioids for chronic pain are critical "to reverse the epidemic of opioid drug overdose deaths and prevent opioid-related morbidity."⁷⁰⁰

- 945. Compounding the harm from deceptive marketing, Defendants worked with Purdue to continue and grow the opioid sales of prescribers that raised red flags of diversion, despite Purdue's legal obligations to report and halt supply. In doing so, it enabled an oversupply of opioids, which allows non-patients to become exposed to opioids, and facilitates access to opioids for both patients who could no longer access or afford prescription opioids and addicts struggling with relapse.
- 946. Most of the illicit use originates from prescribed opioids. It has been estimated that 60% of the opioids that are abused come, directly or indirectly, through physicians' prescriptions.
- 947. As McKinsey itself has recognized in citing a study reaching this conclusion, roughly 80% of heroin users previously used prescription opioids.⁷⁰¹ As many as one in four patients who receive prescription opioids long-term for chronic pain in primary care settings struggles with addiction. And the link between prescription narcotic painkiller abuse and subsequent and/or simultaneous heroin abuse continues to grow.

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⁶⁹⁸ Dart, MD, et al., Trends in Opioid Analgesic Abuse and Mortality in the United States, New Engl. J. Med., 372:241-248 (Jan. 15, 2015).

⁶⁹⁹ Califf, MD, et al., A Proactive Response to Prescription Opioid Abuse, New Engl. J. Med. (Apr. 14, 2016).

⁷⁰⁰ CDC, January 1, 2016 Morbidity and Mortality Weekly Report; Rudd, Rose A., et al. "Increases in drug and opioid overdose deaths - United States, 2000-2014." American Journal of Transplantation 16.4 (2016): 1323-1327.

https://www.mckinsey.com/industries/healthcare-systems-and-services/our-insights/why-we-need-bolder-actionto-combat-the-opioid-epidemic

- 948. In fact, people who are addicted to prescription opioid painkillers are 40 times more likely to be addicted to heroin. The CDC identified addiction to prescription pain medication as the strongest risk factor for heroin addiction. A more recent, and even more deadly problem stemming from the prescription opioid epidemic involves fentanyl, a powerful opioid prescribed for cancer pain or in hospital settings that, in synthetic form, has made its way into Plaintiffs' communities.
- 949. Carfentanil, a powerful derivative of fentanyl, has increasingly been found in heroin and fentanyl sold illicitly. Carfentanil is so strong that it is typically used in veterinary medicine to sedate large wild animals such as elephants, and has been researched as a chemical weapon. A dose the size of a grain of salt can rapidly lead to deadly overdose in humans.
- 950. No demographic is untouched by this epidemic. Nationally, one in five deaths among younger adults in 2016 involved opioids, according to one study. And deaths involving both prescription and illicit opioids have risen sharply, nearly doubling since 2009.
- 951. Opioids were involved in 42% of all fatal drug overdoses in 2015, and another 25% involved heroin. According to the CDC, between 1999 and 2015, more than 183,000 people died in the United States from prescription-related overdoses.
- 952. Rising opioid use and abuse have negative social and economic consequences far beyond overdoses in other respects as well. According to a recent analysis by a Princeton University economist, approximately one out of every three working age men who are not in the labor force take daily prescription pain medication. The same research finds that opioid prescribing alone accounts for 20% of the overall decline in the labor force participation for this group from 2014 to 2016, and 25% of the smaller decline in labor force participation among women. Many of those taking painkillers still said they experienced pain daily.
- 953. In 2009, Dr. Van Zee identified the *precise tactics* that Defendants deployed for all of their opioid clients, including Purdue, as a source of OxyContin misuse and abuse, and suggested

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that regulation may be appropriate to curtail the use of the marketing tactics deployed by Defendants: "The use of prescriber profiling data to target high-opioid prescribers – coupled with very lucrative incentives for sales representatives – would seem to fuel increased prescribing by some physicians – perhaps the most liberal prescribers of opioids and, in some cases, the least discriminate."702

In time, additional evidence mounted supporting the conclusion that Defendants' 954. sales and marketing tactics were demonstrably exacerbating the nationwide opioid crisis. One way of demonstrating the link between aggressive sales and marketing of opioids and worsened mortality outcomes arose out of a quirk of Purdue's own marketing tactics.

In 1996, when OxyContin was introduced, five states – California⁷⁰³, Idaho, Illinois, 955. New York and Texas – maintained "triplicate" programs that required prescribers of Schedule II controlled substances to fill out prescriptions in triplicate.⁷⁰⁴ One of the triplicate copies would then be filed with the state agency in charge of maintaining a prescription database intended to monitor diversion and other potential issues relating to the over-dissemination of Schedule II narcotics. These triplicate programs were precursors to modern prescription drug monitoring programs that have been instituted in nearly every state in response to the opioid crisis.

956. Purdue recognized that the requirement to submit records of controlled substance prescriptions to a governmental database chilled prescribers' willingness to prescribe medications subject to the constraints of the triplicate programs. Because Purdue viewed these triplicate

⁷⁰² Art Van Zee, The Promotion and Marketing of OxyContin: Commercial Triumph, Public Health Tragedy, 99 AM. PUB. **HEALTH** 221. 221. (Feb. 2009), available at: https://www.ncbi.nlm.nih.gov/pmc/articles/PMC2622774/pdf/221.pdf.

⁷⁰³ California was the first state to implement a triplicate program in response to concerns about the diversion of opiumbased pharmaceuticals. The year was 1939. See Abby E. Alpert, William N. Evans, Ethan M.J. Lieber, and David Powell, Origins of the Opioid Crisis and its Enduring Impacts, NBER Working Paper No. 26500, November 2019, available at: https://www.nber.org/papers/w26500; See also, supra., fn. 1.

requirements as an overly burdensome hindrance on prescribing, the company chose to focus its marketing efforts in other states that did not impose these constraints.

957. This resource-allocation decision by Purdue to focus more marketing efforts in states with fewer regulations regarding the prescribing of controlled substances provided a way to test whether *marketing* of OxyContin, by itself, was a cause of not only increased overdose rates for OxyContin, but of *all* opioid-related overdoses, *including* those involving illicit opioids such as heroin and fentanyl.

958. The results were stark. In 2019, economists from the University of Pennsylvania, Notre Dame, and the RAND Corporation analyzed the disparate outcomes in overall opioid overdose mortality experienced in the triplicate states, where Purdue did not focus its marketing efforts, and non-triplicate states where Purdue did focus those efforts.⁷⁰⁵

959. The economists found that "OxyContin distribution was about 50% lower in 'triplicate states' in the years after the launch. While triplicate states had higher rates of overdose deaths prior to 1996, this relationship flipped shortly after the launch [of OxyContin] and triplicate states saw substantially slower growth in overdose deaths, continuing even twenty years after OxyContin's introduction. Our results show that the introduction and marketing of OxyContin explain a substantial share of overdose deaths over the last two decades."⁷⁰⁶

960. A 2017 *Journal of American Medical Association* study found that physicians ordered fewer promoted brand-name medications and prescribed more cost-effective generic versions if they worked in hospitals that instituted rules about when and how pharmaceutical sales representatives were allowed to detail prescribers.⁷⁰⁷ The changes in prescribing behavior appeared

⁷⁰⁵ Abby E. Alpert, William N. Evans, Ethan M.J. Lieber, and David Powell, *Origins of the Opioid Crisis and its Enduring Impacts*, NBER Working Paper No. 26500, November 2019, *available at*: https://www.nber.org/papers/w26500

⁷⁰⁶ *Id.* (emphasis added).

⁷⁰⁷ Ian Larkin et al., Association Between Academic Medical Center Pharmaceutical Detailing Policies and Physician Prescribing, 317 J. Am. Med. Ass'n 1785 (2017).

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strongest at hospitals that implemented the strictest detailing policies and included enforcement measures. Another study involved the research of four different practices which included visits by sales representatives, medical journal advertisements, direct-to-consumer advertising, and pricing, and found that sales representatives have the strongest effect on driving drug utilization. An additional study found that doctor meetings with sales representatives are related to changes in doctor prescribing practices and requests by physicians to add the drugs to hospitals' formularies.

- 961. Separately, a recent Journal of American Medical Association study analyzed the Centers for Medicare and Medicaid Services' Open Payments database regarding pharmaceutical company marketing efforts towards doctors, as well as CDC data on prescription opioid overdose deaths and prescribing rates, in order to assess whether pharmaceutical marketing of opioids to physicians affected the rate of prescription opioid overdose deaths. Notably, the study analyzed these marketing practices beginning August 1, 2013, and ending December 31, 2015.⁷⁰⁸
- 962. Those dates are significant, as the study captures the same timeframe that McKinsey's Project Turbocharge, re-christened *E2E*, was implemented.
- 963. The study noted "physician prescribers are the most frequent source of prescription opioids for individuals who use opioids nonmedically."⁷⁰⁹
- 964. The study found that "increased county-level opioid marketing was associated with elevated overdose mortality 1 year later, an association mediated by opioid prescribing rates; per capita, the number of marketing interactions with physicians demonstrated a stronger association with mortality than the dollar value of marketing."⁷¹⁰

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⁷⁰⁸ Scott E. Hadland et. al., Association of Pharmaceutical Industry Marketing of Opioid Products with Mortality from 27 Opioid-Related Overdoses, JAMA Network. January https://jamanetwork.com/journals/jamanetworkopen/fullarticle/2720914.

²⁸ ⁷⁰⁹ *Id*.

⁷¹⁰ <u>Id.</u> (emphasis added)

965. Referring to the types of sales and marketing tactics Publicis provided to its clients, and helped them implement, the authors concluded, "amid a worsening opioid crisis, our results suggest that industry marketing to physicians may run counter to current efforts to curb excessive opioid prescribing."⁷¹¹

966. The authors' proposed solution was plain simple, and echoed Dr. Van Zee's congressional testimony from 2002: "Pharmaceutical companies might also consider, as one manufacturer recently did, voluntarily ceasing marketing opioid products directly to physicians." 712

967. Incredibly, in an August 7, 2016, presentation to Purdue regarding their "Corporate Identity Transformation," Publicis offered, as one strategy *option* to be *considered* among others, voluntarily ceasing to market opioids in order to "fully embrace a deeper-held responsibility for progress in pain and keeping people safe." The pain franchise could be placed "on probation." It would be a "bold commitment:"

⁷¹¹ *Id*. ⁷¹² *Id*.

incentivized to sell products,

Purdue can commit to more

responsibly serving physicians

by employing only an

expanded team of Medical

Science Liaisons (MSLs) for

managing 1-to-1 personal

interactions with HCPs

PUT THE PAIN FRANCHISE "ON PROBATION" ACTIONS THAT DEMONSTRATE A BOLD COMMITMENT Take clear, bold, and direct actions at the corporate level to show the public that Purdue is willing to compromise its own business in order to fully embrace a deeper-held responsibility for progress in pain and keeping people safe Shut Down the Sales Force (Shift to MSL Model) Achieve Full HCP Compliance with ER-LA REMS Program While the sales force is Aspire to regulate who Boldly commit to the goal

Aspire to regulate who is qualified to prescribe opioids. Purdue leads the effort working with FDA and other pharma companies to up its commitment to getting HCPs REMS-compliant it They're going to prescribe opioid pain meds.

Boldly commit to the goal of getting every patient att therapy: employ systems and strategies across all brands to help HCPs and Patients better manage their pain regimen and plan for the eventual discontinuation of pain therapy.

razorfish health

968. Defendants' role in the opioid industry was central. Given its overlapping engagements with multiple opioid manufacturers regarding the sales of competing products **at the same time**, Defendants were in a position to know the intricacies of the sales and marketing tactics of competing opioid sellers, including confidential information and trade secrets of these clients. And Publicis served as matchmaker to Defendant Practice Fusion, ushering it in to service opioid accounts like Purdue's that Publicis already controlled.

969. All the while, Defendants' clients were repeatedly subjected to enforcement actions for their work selling opioids both before and during the pendency of Defendants' client relationships. For instance, after its 2007 guilty plea with the United States Department of Justice ("DOJ"), Purdue Pharma settled with the State of Kentucky in 2015 for \$24 million.⁷¹³ The

⁷¹³ See the December 22, 2015, settlement between the Commonwealth of Kentucky and Purdue Pharma Inc., available at https://ag.ky.gov/pdf news/purduepharmaoxycontin.pdf.

settlement concerned similar conduct as the 2007 guilty plea, including the sales and marketing of Purdue's opioids. Defendants were all active participants in and contributors to this conduct.

- 970. Because of their extensive and long-term client relationships, Defendants were in a unique position to know how the entire industry's opioid sales and marketing tactics were playing out, both in terms of return on investment for their individual clients, as well as overall market trends such as the rise of the opioid crisis. Endo may not have known the specifics of competitor Purdue's marketing efforts for OxyContin, just as Purdue may not have known the specifics of Janssen's Nucynta plan. But McKinsey, Publicis, and ZS knew all three, as well as what Teva and others were doing with their own branded opioid sales and marketing efforts *in real time*.
- 971. As the modern opioid epidemic became apparent and the subject of nationwide attention, Defendants toiled diligently behind the scenes of several opioid manufacturers in the pursuit of one goal: maximizing volumes and profits from the sale of these addictive and deadly Schedule II controlled substances.
- 972. And so, through the affirmative acts undertaken by the Defendants, history repeats itself. This time, however, the devastation is on a scale previously unimaginable.

V. TOLLING OF STATUTES OF LIMITATION

- 973. Defendants are equitably estopped from relying upon a statute of limitations defense. Alongside their clients, Defendants undertook active efforts to deceive Plaintiff and to purposefully conceal its unlawful conduct and fraudulently assure the public, including Plaintiff, that opioids were non-addictive, effective, and safe for the treatment of long-term chronic pain and non-acute, non-cancer pain with the goal of increased sales, greater availability and access to opioids, and maximizing profits.
- 974. Defendants and their clients were deliberate in taking steps to conceal their conspiratorial behavior and active role in the deceptive marketing of opioids. This deceptive

marketing—which included the above falsehoods that opioids were safer, less subject to abuse, and less addictive than other pain medications—was a substantial factor in the oversupply of opioids through overprescribing and suspicious sales, all of which fueled the opioid epidemic.

975. Defendants deliberately advised their clients on marketing strategies and tactics to bolster their opioid products as non-addictive, safe, and efficacious without reliable scientific evidence to support same, and implemented the same. Defendants' services were given confidentially, and both Defendants and their clients concealed the content of those services from the public. In doing so, Defendants concealed their role in shaping, editing, and providing the content of the false and misleading materials addressing pain management and opioids that were widely disseminated to regulators, prescribers, and the public at large, including Plaintiff.

976. Defendants also concealed from Plaintiff the existence of the Plaintiff's claims by hiding their and their client's lack of cooperation with law enforcement. For example, in May 2007, Purdue and three of its executives pled guilty to federal charges of misbranding OxyContin in what the company acknowledged was an attempt to mislead doctors about the risk of addiction and entered into a Corporate Integrity Agreement explained above. Purdue was ordered to pay \$600 million in fines and fees. In its plea, Purdue admitted that its promotion of OxyContin was misleading and inaccurate, misrepresented the risk of addiction, and was unsupported by science. Additionally, Michael Friedman, the company's president, pled guilty to a misbranding charge and agreed to pay \$19 million in fines; Howard R. Udell, Purdue's top lawyer, also pled guilty and agreed to pay \$8 million in fines; and Paul D. Goldenheim, its former medical director, pled guilty as well and agreed to pay \$7.5 million in fines.

977. Nevertheless, even after the guilty pleas, Purdue continued to pay doctors on speakers' bureaus to promote the liberal prescribing of OxyContin for chronic pain and fund seemingly neutral organizations to disseminate the message that opioids were non-addictive as well

as other misrepresentations. Purdue also assembled an army of lobbyists to fight any legislative actions that might encroach on its business. Between 2006 and 2015, Purdue and other painkiller producers, along with their associated nonprofits, spent nearly \$900 million dollars on lobbying and political contributions—eight times what the gun lobby spent during that period. Defendants participated extensively in these actions and provided Purdue with strategies and assistance to maximize sales as described in this Complaint. Defendants knew that the actions they took with Purdue were unlawful, and yet deliberately proceeded in order to increase Purdue's sales and profits, and in turn to serve Defendants' financial interests.

978. Defendants affirmatively sought to convince the public that its clients' legal duties to report suspicious sales of opioids had been satisfied through public assurances that they were working to curb the opioid epidemic. For example, after the 2007 Purdue guilty plea described above, Defendants provided services to protect the company's public image and sales, aiding in the concealment of the addictive nature and dangers associated with opioid use and denying blame for the epidemic, attributing it instead solely to abuse and inappropriate prescribing. At the guidance and advice of Defendants, Purdue and others publicly portrayed themselves as committed to working diligently with law enforcement and others to prevent diversion of these dangerous drugs and curb the opioid epidemic, and they made broad promises to change their ways, insisting they were good corporate citizens. Instead, Defendants assisted Purdue, for example, with marketing campaigns and messaging that continued business as usual, indiscriminately targeting high prescribers and promoting opioids as safe but avoiding the pitfalls of the Corporate Integrity Agreement. These repeated misrepresentations misled regulators, prescribers, and the public, including the Plaintiff, and deprived Plaintiff of actual or implied knowledge of facts sufficient to put the Plaintiff on notice of potential claims.

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979. Plaintiff did not discover the nature, scope, and magnitude of Defendants' misconduct, and its full impact on Plaintiff, and could not have acquired such knowledge earlier through the exercise of reasonable diligence.

- 980. Prior to the applicable limitations period, Plaintiff did not suspect, and had no reason to suspect, that Defendants' conduct caused their injuries, including the consumption of Plaintiff's resources as the opioid epidemic remains unabated.
- 981. Defendants intended that its actions and omissions made with its clients would be relied upon, including by the Plaintiff. The Plaintiff did not know and did not have the means to know the truth due to Defendants' and their clients' actions and omissions.
- 982. The Plaintiff reasonably relied on the affirmative statements developed by Defendants and made by their clients regarding their purported compliance with their obligations under the law and consent orders, which were false and only intended to save the clients' public image.
- 983. Defendants' fraudulent concealment has tolled the running of any statute of limitations. Through their and their clients' affirmative misrepresentations and omissions, Defendants actively concealed from Plaintiff the risks associated with opioids that led to the opioid crisis. The wrongdoing, misrepresentations, and omissions by Defendants have not ceased because the public nuisance remains unabated.

VI. CAUSES OF ACTION

COUNT I:Racketeer Influenced and Corrupt Organizations Act (RICO)

984. Plaintiffs re-allege all of the foregoing allegations and incorporate them herein by reference.

- 985. This claim is brought by Plaintiffs against Defendants for actual damages, treble damages, and available injunctive and/or equitable relief under 18. U.S.C. § 1964, for violations of 18 U.S.C. § 1961, *et seq.*, specifically, 18 U.S.C. § 1962(c) and (d).
- 986. Section 1962(c) makes it "unlawful for any person employed by or associated with any enterprise engaged in, or the activities of which affect, interstate or foreign commerce, to conduct or participate, directly or indirectly, in the conduct of such enterprise's affairs through a pattern of racketeering activity" 18 U.S.C. § 1962(c).
- 987. At all relevant times, each Defendant is and has been a "person" under 18 U.S.C. § 1961(3) because each is capable of holding, and does hold, "a legal or beneficial interest in property."
- 988. Plaintiffs are each a "person," as the term is defined in 18 U.S.C. § 1961(3), and have standing to sue under 18 U.S.C. § 1964(c) as they were and are injured in their business and/or property "by reason of" the RICO Act violations described herein.
- 989. Section 1962(d) makes it unlawful for "any person to conspire to violate" Section 1962(c), among other provisions. See 18 U.S.C. § 1962(d).
- 990. Defendants conducted the affairs of an enterprise through a pattern of racketeering activity, in violation of 18 U.S.C. § 1962(c) and § 1962(d).

Description of the Enterprise

991. Section 1961(4) defines an enterprise as "any individual, partnership, corporation, association, or other legal entity, and any union or group of individuals associated in fact although not a legal entity." 18 U.S.C. § 1961(4).

- 992. Under 18 U.S.C. § 1961(4), a RICO "enterprise" may be an association-in-fact that, although it has no formal legal structure, has (i) a common purpose, (ii) relationships among those associated with the enterprise, and (iii) longevity sufficient to pursue the enterprise's purpose. *See Boyle v. United States*, 556 U.S. 938, 946 (2009).
- 993. Opioid manufacturers, including Purdue, Johnson & Johnson, Janssen, Cephalon, Endo, and Mallinckrodt (collectively the "Opioid Manufacturers"), together with Defendants and McKinsey ("the Opioid Consultants"), participated in the marketing and sale of opioids as described in this Complaint, (collectively, the "Opioid Marketing Enterprise Members" or the "Enterprise Members") engaged in a scheme to unlawfully increase sales of opioids— both by growing their share of the prescription painkiller market *and* by growing the market as a whole—through repeated and systematic misrepresentations, concealments, and omissions of material fact about the safety and efficacy of opioids for treating long-term chronic pain, together with other deceptive and fraudulent acts and practices, as described in the Factual Allegations section of this Complaint and in the Master Consolidated Complaint filed by Plaintiffs in MDL 2996 on December 6, 2021, *cf.* Dkt. 300 (Tribal Plaintiffs Master Complaint).
- 994. In order to unlawfully increase the demand for opioids and thereby increase their own profits despite their knowledge of the harmful effects that would follow, the Opioid Marketing Enterprise Members formed an association-in-fact enterprise (the "Opioid Marketing Enterprise" or the "Enterprise"). The Opioids Manufacturers worked together to accomplish their aims, with McKinsey and Defendants serving as go-betweens that held all of the companies together and helped design and coordinate the deceptive marketing and sales strategies. Through McKinsey and

Defendants and their own personal relationships, the members of the Opioid Marketing Enterprise had the opportunity to form and take actions in furtherance of the Opioid Marketing Enterprise's common purpose: lying to prescribers and Plaintiffs in order to increase sales of addictive and dangerous drugs and line the enterprise members' pockets. The Opioid Marketing Enterprise Members' substantial financial contributions to the Opioid Marketing Enterprise and the advancement of opioids-friendly messaging fueled the U.S. opioid epidemic.

995. In the alternative, the association-in-fact Opioid Marketing Enterprise existed just between McKinsey, Defendants, and Purdue, who worked together to unlawfully increase sales of *all* opioids—not just Purdue's own products—through repeated and systematic misrepresentations about the safety and efficacy of opioids for treating long-term chronic pain. McKinsey and Defendants knew Purdue was marketing its opioids illegally and fueling an opioid epidemic, but using the knowledge gained from Defendants' and McKinsey's work with other opioid manufacturers, Defendants and McKinsey joined forces with Purdue to turbocharge the opioids market in order to profit from this crisis.

996. The Controlled Substances Act (the "CSA") and its implementing regulations require that "[e]very person who manufactures, distributes, dispenses, imports, or exports any controlled substance," including opioids, become a "registrant." *See* 21 U.S.C. § 823(a)-(b); 21 C.F.R. § 1301.11(a). These registrants, including opioid manufacturer and distributors, must maintain a system to identify and report suspicious orders, including orders of unusual size or frequency, or orders deviating from a normal pattern, and maintain effective controls against diversion of controlled substances. *See* 21 U.S.C. § 823; 21 C.F.R. § 1301.74(b).

997. Despite these duties, Defendants, McKinsey and the other Enterprise Members engaged in a scheme with the overarching purpose of materially expanding prescription opioid use by altering the medical community's opioid prescribing practices through repeated fraudulent

statements and misrepresentations. The Opioid Marketing Enterprise's scheme was sophisticated, well-developed, and fraudulent and was designed to increase the prescription rate for opioid medications the Enterprise Members knew where dangerous and highly addictive. At all relevant times, Defendants and McKinsey were aware of the conduct of the Enterprise, were knowing and willing participants in that conduct, and reaped profits from that conduct in the form of payments from other Enterprise Members as a reward for work done to increase sales and distribution of prescription opioids.

The Common Purpose and Scheme of the Opioid Marketing Enterprise.

998. The Opioid Marketing Enterprise Members, through the Opioid Marketing Enterprise, concealed the true risks and dangers of opioids from the medical community and Plaintiffs and made misleading statements and misrepresentations about opioids that downplayed the risk of addiction and exaggerated the benefits of opioid use. These misleading statements included: (1) that addiction is rare among patients taking opioids for pain; (2) that addiction risk can be effectively managed; (3) that symptoms of addiction exhibited by opioid patients are actually symptoms of an invented condition, which the Opioid Marketing Enterprise Members named "pseudoaddiction"; (4) that withdrawal is easily managed; (5) that increased dosing presents no significant risks; (6) that long-term use of opioids improves function; (7) that the risks of alternative forms of pain treatment are greater than the adverse effects of opioids; (8) that use of time-released dosing prevents addiction; and (9) that abuse-deterrent formulations provide a solution to opioid abuse.

999. The scheme devised, implemented, and conducted by the Opioid Marketing Enterprise Members was a common course of conduct designed to ensure that the Opioid Marketing Enterprise Members unlawfully increased their sales and profits through concealment and misrepresentations about the addictive nature and effective use of the Opioid Manufacturers' drugs.

The Opioid Marketing Enterprise Members acted together for a common purpose and perpetuated the Opioid Marketing Enterprise's scheme.

1000. There was regular communication between the Opioid Marketing Enterprise Members in which information was shared, misrepresentations were coordinated, and payments were exchanged. The Opioid Marketing Enterprise Members functioned as a continuing unit for the purpose of implementing the Opioid Marketing Enterprise's scheme and common purpose, and each agreed and took actions to hide the scheme and continue its existence.

1001. As public scrutiny and media coverage focused on how opioids ravaged communities throughout the United States, Defendants and McKinsey did not challenge Purdue or other manufacturers' misrepresentations, seek to correct their previous misrepresentations, terminate their role in the Opioid Marketing Enterprise, nor disclose publicly that the risks of using opioids for chronic pain outweighed their benefits and were not supported by medically acceptable evidence. Instead, despite its knowledge of the ongoing fraud and the danger it posed, Defendants and McKinsey continued to participate in the Opioid Marketing Enterprise for financial gain.

1002. The impact of the Opioid Marketing Enterprise's scheme is still in place—i.e., the opioids continue to be prescribed and used for chronic pain throughout the United States, and the epidemic continues to injure Plaintiffs and consume the resources of Plaintiffs.

1003. The evidence shows that the Opioid Marketing Enterprise Members, including Defendants and McKinsey, were each willing participants in the Opioid Marketing Enterprise, had a common purpose and interest in the object of the scheme, and functioned within a structure designed to effectuate the Enterprise's purpose.

The Conduct of the Opioid Marketing Enterprise Violated Civil RICO.

1004. From at least 2004 to the present, each of the Opioid Marketing Enterprise Members played some part in directing the affairs of the Opioid Marketing Enterprise and participated in the operation or management of the affairs of the Opioid Marketing Enterprise, directly or indirectly, in the following ways:

1005. Creating and providing a body of deceptive, misleading, and unsupported medical and popular literature about opioids that (i) understated the risks and overstated the benefits of long-term use; (ii) appeared to be the result of independent, objective research; and (iii) was thus more likely to be relied upon by physicians, patients, and payors;

1006. Creating and providing a body of deceptive, misleading, and unsupported electronic and print advertisements about opioids that (i) understated the risks and overstated the benefits of long-term use; (ii) appeared to be the result of independent, objective research; and (iii) was thus more likely to be relied upon by physicians, patients, and payors;

1007. Creating and providing a body of deceptive, misleading, and unsupported sales and promotional training materials about opioids that (i) understated the risks and overstated the benefits of long-term use; (ii) appeared to be the result of independent, objective research; and (iii) was thus more likely to be relied upon by physicians, patients, and payors;

1008. Devising and implementing marketing schemes that included targeting and misleading physicians, unlawfully incentivizing sales representatives to maximize prescriptions and dosages, and evading regulatory constraints; and

1009. Disseminating many of their false, misleading, imbalanced, and unsupported statements through unbranded materials that appeared to be independent publications.

1010. The scheme devised and implemented by the Opioid Marketing Enterprise Members amounted to a common course of conduct intended to enrich themselves by increasing sales of prescription opioids by convincing doctors to prescribe and patients to use opioids, including for

long-term chronic pain, despite the Opioid Marketing Enterprise Members' knowledge of the addictions and deaths that would occur as a result. The scheme was a continuing course of conduct, and many aspects of it continue through to the present.

The Opioid Marketing Enterprise Members Conducted or Participated, Directly or Indirectly, in the Conduct of the Enterprise's Affairs.

1011. "[T]o conduct or participate, directly or indirectly, in the conduct" of an enterprise, "one must participate in the operation or management of the enterprise itself." *Reves v. Ernst & Young*, 507 U.S. 170, 185 (1993).

1012. As described herein, the Opioid Marketing Enterprise Members participated in the conduct of the Enterprise through a pattern of racketeering activity, and Defendants and McKinsey were the masterminds of marketing schemes deployed by the Enterprise members to defraud prescribers and Plaintiffs by using the mail and wires in furtherance of plans that were designed with specific intent to defraud.

1013. The Opioid Marketing Enterprise Members conducted an association-in-fact enterprise and/or participated in the conduct of an enterprise through a pattern of illegal activities (the predicate racketeering acts of mail and wire fraud) to carry-out the common purpose of the Opioid Marketing Enterprise, i.e., to unlawfully increase profits and revenues from the continued prescription and use of opioids for long-term, chronic pain. Through the racketeering activities of the Opioid Marketing Enterprise, the Opioid Marketing Enterprise Members sought to further the common purpose of the Enterprise through a fraudulent scheme to change prescriber habits and public perception about the safety and efficacy of opioid use. In so doing, each of the Opioid Marketing Enterprise Members knowingly conducted and participated in the conduct of the Enterprise by engaging in mail and wire fraud, in violation of 18 U.S.C. §§ 1962(c) and (d).

1014. The Opioid Marketing Enterprise is an association-in-fact enterprise that consists of the Opioid Marketing Enterprise Members.

1015. Each of the Opioid Marketing Enterprise Members conducted and participated in the conduct of the Opioid Marketing Enterprise by playing a distinct role in furthering the Enterprise's common purpose of increasing profits and sales through the knowing and intentional dissemination of false and misleading information about the safety and efficacy of long-term opioid use, and the risks and symptoms of addiction, in order to increase the market for prescription opioids by changing prescriber habits and public perceptions.

1016. Specifically, the Opioid Marketing Enterprise Members each worked together to coordinate the Enterprise's goals and conceal their role, and the Enterprise's existence, from prescribers and Plaintiffs by, among other things, (i) funding, creating, editing, and distributing publications that supported and advanced their false messages; (ii) funding key opinion leaders ("KOLs") to further promote their false messages; and (iii) tasking their own employees to direct deceptive marketing materials and pitches directly at physicians.

1017. Further, each of the Opioid Marketing Enterprise Members had systematic links to, and personal relationships with, each other through joint participation in lobbying groups, trade industry organizations, contractual relationships, and continuing coordination of activities. The systematic links and personal relationships that were formed and developed allowed the Opioid Marketing Enterprise Members the opportunity to form the common purpose and agree to conduct and participate in the conduct of the Opioid Marketing Enterprise. Specifically, each of the Opioid Marketing Enterprise Members coordinated their efforts through the same KOLs and front groups, based on their agreement and understanding that the front groups and KOLs were industry friendly and would work together with the Opioid Marketing Enterprise Members to advance the common purpose of the Opioid Marketing Enterprise; and each of the individuals and entities who formed

the Opioid Marketing Enterprise acted to enable the common purpose and fraudulent scheme of the Opioid Marketing Enterprise.

1018. At all relevant times, the Opioid Marketing Enterprise: (a) had an existence separate and distinct from each Opioid Manufacturer and its members; (b) was separate and distinct from the pattern of racketeering in which the Opioid Marketing Enterprise Members engaged; (c) was an ongoing and continuing organization consisting of individuals, persons, and legal entities, including each of the Opioid Marketing Enterprise Members; (d) was characterized by interpersonal relationships between and among each member of the Opioid Marketing Enterprise; and (e) had sufficient longevity for the Enterprise to pursue its purpose and functioned as a continuing unit.

1019. The Opioid Marketing Enterprise Members conducted and participated in the conduct of the Opioid Marketing Enterprise through a pattern of racketeering activity that employed the use of mail and wire facilities, in violation of 18 U.S.C. § 1341 (mail fraud) and § 1343 (wire fraud), to increase profits and revenue by changing prescriber habits and public perceptions in order to increase the prescription and use of prescription opioids and expand the market for opioids.

1020. The Opioid Marketing Enterprise Members each committed, conspired to commit, and/or aided and abetted in the commission of at least two predicate acts of racketeering activity (i.e., violations of 18 U.S.C. §§ 1341 and 1343) within the past ten years. The multiple acts of racketeering activity that the Opioid Marketing Enterprise Members committed, or aided and abetted in the commission of, were related to each other, posed a threat of continued racketeering activity, and therefore constitute a "pattern of racketeering activity." The racketeering activity was made possible by the Opioid Marketing Enterprise Members' regular use of the facilities, services, distribution channels, and employees of the Opioid Marketing Enterprise, the U.S. Mail, and interstate wire facilities. The Opioid Marketing Enterprise Members participated in the scheme to

defraud by using mail, telephones, and the internet to transmit communications and payments in interstate or foreign commerce.

The Conduct was More than a Typical Business Relationship.

1021. There were strong relationships among those associated with the Opioid Enterprise and sufficient longevity among Enterprise associates to pursue the Enterprise's common purpose. The common purpose was to increase opioid revenues unlawfully by misrepresenting and lying about opioids in order to changing prescriber habits and the perception regarding the safety and efficacy of opioids for chronic pain and long-term use. The Enterprise's deceit was, in part, in its failure to disclose that increasing strength and dosing actually increased the risk of addiction and overdose and that patients on opioids for more than a brief period develop tolerance, requiring increasingly high doses to achieve pain relief.

1022. On March 1, 2004, McKinsey entered into a "Master Consulting Agreement" with Purdue for "services that would be defined from time to time."⁷¹⁴ The Master Consulting Agreement was signed by then-McKinsey director Rob Rosiello."⁷¹⁵

1023. From 2004 through 2008, McKinsey advised Purdue on research and development, business development, and product licensing related to Purdue's opioid products.⁷¹⁶ Consistent with its business model, McKinsey leveraged these projects into growth of its "Broader Strategy work" also underway with Purdue.⁷¹⁷ Specifically, in October 2008, Purdue retained McKinsey for broad strategy work after two board members "blessed" Purdue executive Craig Landau with doing "whatever he thinks is necessary to 'save the business" after the 2007 criminal plea and introduction of generic competition to the older OxyContin.⁷¹⁸ Purdue relied heavily on McKinsey

⁷¹⁴ MCK-MDL2996-0085849; PPLPC012000069192

⁷¹⁵ MCK-MDL2996-0085849, at 0085880.

⁷¹⁶ PPLPC013000116218; PPLP004401340

⁷¹⁷ MCK-MAAG-0117875

⁷¹⁸ MCK-MAAG-0117875

to help Purdue publicly portray itself as a good corporate citizen who could now be trusted and was even working on an "abuse-deterrent" or "ADF" form of OxyContin. Defendant Publicis worked intimately with McKinsey is designing and disseminating these messages to their intended audience.

1024. Over their many years of working together, McKinsey and Richard Sackler developed a close relationship. Indeed, one McKinsey partner, Maria Gordian, describes herself as a counselor to Richard Sackler in an "Ey 2009 Impact Summary."⁷¹⁹

1025. For Publicis' part, they owned their client. Publicis' Karl Tiedemann noted in an email the "amazing relationship" developing between Purdue and Publicis, and quoted a Purdue employee as stating that the Hysinglia account was "the final piece and we now own Purdue." (emphasis added).

Rather, the members of the Enterprise knew that opioids were addictive and causing serious harm to people and communities but chose to work together to lie to prescribers and Plaintiffs about these drugs in order to increase their bottom lines. Defendants and McKinsey worked closely with the Opioid Manufacturers to achieve these aims. Defendants and McKinsey, as advisors to multiple Opioid Manufacturers, also had access to information about multiple players and was able to coordinate the fraud occurring across the Enterprise. As discussed below, McKinsey was particularly embedded in Purdue's organizational structure and the relationship's longevity was sufficient to pursue the Enterprise's purposes. During the 2009-2014 period in particular, Purdue relied extensively on McKinsey to develop its sales and marketing strategy for OxyContin. Publicis worked with McKinsey and Purdue to craft and disseminate these strategies by using targeting and

⁷¹⁹ MCK-MAAG-0118669

28 721 MCK-MAAG-0118669 722 PPLPC061000045395

salesforce optimization methods designed by ZS and distribution channels offered by Practice Fusion.

strengthen their relationships with Purdue and assist Purdue in selling opioids after Purdue's 2007 criminal guilty plea. As part of the guilty plea, Purdue admitted that its "supervisors and employees, with the intent to defraud or mislead, marketed and promoted OxyContin as less addictive, less subject to abuse and diversion, and less likely to cause tolerance and withdrawal than other pain medication." But rather than be deterred by this, Defendants and McKinsey dove in. In a March 2009 self-assessment, Ms. Gordian described McKinsey's progress in having "continue[d] to expand the depth and breadth of [its] relationships with Purdue" and plans to "deepen[]" McKinsey's "relationship with the Sackler family," including by "serving them on key business development issues" and "expanding" McKinsey's relationship with members of Purdue's senior management team. Meanwhile, internal Rosetta communications refer to Publicis "owning" Purdue given the extent of their relationship with the client, to wit, "we now own them."

1028. By August 2009, Richard Sackler had convened a meeting of Purdue board members and staff to discuss efforts to "reverse the decline in the OxyContin tablets market." During the 2009-2014 period in particular, Purdue relied extensively on Defendants and McKinsey to develop and implement sales and marketing strategy for OxyContin. Defendants and McKinsey worked closely with Purdue on both the creation and implementation of OxyContin sales strategy. McKinsey's work for Purdue included consulting, review of product acquisition, evaluation of research and development, advising Purdue on the design of clinical studies, risk management, and product marketing. 723

⁷²⁰ Information at pp. 5-6, *United States v. Purdue Frederick Co.*, No. 07-cr-29-JPJ (W.D. Va. May 10, 2007), Doc. 5.

⁷²² PPLPC061000045395 ⁷²³ PPLPC029000547371

1029. On May 28, 2013, McKinsey entered into a "Statement of Services to the Master Consulting Agreement" (the "2013 Agreement") with Purdue to "conduct a rapid assessment of the underlying drivers of current OxyContin performance, identify key opportunities to increase nearterm OxyContin revenue and develop plans to capture priority opportunities."⁷²⁴ The 2013 Agreement stated, "We have a long history of partnership with Purdue, and we would make best efforts to leverage our understanding of your business—both in terms of content and culture." The 2013 Agreement was signed by then-principal Arnab Ghatak who would "lead the team with senior leadership from Rob Rosiello and Martin Elling."

1030.

Throughout that time period, Defendants worked alongside Purdue and McKinsey to design, implement and disseminate these strategies.

1031. Thereby, even after the 2007 guilty plea, Purdue, with Defendants and McKinsey's ongoing aid and assistance, saw growing profits from opioid sales. In 2015 alone, Purdue obtained \$3 billion in annual opioid sales—a four-fold increase from its 2006 sales of \$800 million.

1032. Defendants and McKinsey's relationship with Purdue went far beyond a typical business relationship. Defendants and McKinsey worked closely with Purdue on both the creation and implementation of OxyContin sales strategy, a strategy Defendants and McKinsey knew had been based on misleading and defrauding doctors and patients alike about a dangerous and highly addictive drug. McKinsey and Defendants interacted with Purdue at all levels of the corporate

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⁷²⁴ Excerpt from U.S. Department of Justice Plea Agreement with Purdue Pharma L.P. October 20, 2020. 18, ¶88. https://www.justice.gov/opa/press-release/file/1329576/download.

⁷²⁵ PPLPC018001462324

hierarchy – from fields sales representative all the way up to Purdue's Sackler-controlled board of directors.

1033. Further, Defendants and McKinsey had access to detailed prescribing information enabling them to determine if there were suspicious or problematic prescribing patterns. Rather than using this information to help their clients prevent diversion of controlled substances, Defendants and McKinsey and the Opioid Marketing Enterprise used this information in furtherance of their scheme to defraud prescribers and Plaintiffs, target and increase sales to prescribers who were overprescribing, and continue to fuel opioid addiction and the resulting epidemic.

The Fraudulent Schemes

1034. As detailed above, the operation of the Opioid Marketing Enterprise, included several schemes to defraud that helped to further the goals its members—i.e., to expand the market and increase profits and sales through the knowing and intentional dissemination of false and misleading information about the safety and efficacy of long-term opioid use, and to increase profits for the Enterprise Members via expanding the market for opioids.

Fraudulent Marketing Scheme: Deceptive Messaging Regarding Opioid Use

1035. As described throughout, Defendants and McKinsey sought to unlawfully increase profits and revenues from the continued prescription and use of opioids for long-term, chronic pain by changing prescriber habits and public perception regarding the safety and efficacy of opioids. Defendants' and McKinsey's fraud specifically targeted prescribers and set out to convince them that they should prescribe more and more opioids, overcoming what could otherwise be a check on opioid manufacturers ability to increase sales of addictive products.

1036. Despite Defendants and McKinsey knowing that reformulated OxyContin could still be abused, having advised Purdue on the design of tests of reformulated OxyContin as part of

Purdue's FDA submission,⁷²⁶ in furtherance of the scheme to defraud, Defendants and McKinsey spread messages that prescribing opioids could provide "freedom" and "peace of mind" for its users and that physicians could "tailor the dose."

1037. After Purdue's 2007 criminal plea for illegally marketing OxyContin, Defendants and McKinsey created strategies to repair Purdue's reputation and boost OxyContin sales. In 2008, Purdue submitted a New Drug Application for a reformulation of OxyContin, ostensibly to make it more difficult to abuse by extracting the active ingredient from it or otherwise defeating the time-release mechanism in OxyContin tablets—i.e., another product Purdue would later deceptively promote as safer than and less prone to abuse than it was.

1038. In June 2009, McKinsey helped Purdue prepare for an FDA advisory committee meeting.

OxyContin: Question & Answer Book" in September 2009, with questions including "Why should we trust you?" In response, McKinsey recommended Purdue say "We acknowledge mistakes made in the past[;]" "We have x, y and z measures in place that did not exist before[;]" and "[a]t all levels, Purdue's focus is on maintaining the highest ethical standards and meeting the needs of patients[.]" To the question of "Who at Purdue takes personal responsibility for all these deaths?[,]" McKinsey recommended Purdue say, "We all feel responsible[.]"

⁷²⁶ McK-MAAG-0118669

727 PDD8901645845

Id

⁷²⁹ MCK-MAAK-0152135

1040. As described above, Defendant Publicis also engaged in work with the FDA, and through that work was able to disseminate messages consistent with Purdue's strategy to increase overall opioid prescribing.

1041. Defendants and McKinsey and the other Opioid Marketing Enterprise Members knew the changes Purdue made would not make opioids non-addictive or prevent them from being used to create and further substance abuse problems. For example, in 2009, the FDA noted in permitting ADF labeling that "the tamper-resistant properties will have no effect on abuse by the oral route (the most common mode of abuse)." Similarly, in approving reformulated OxyContin, the FDA cautioned that the reformulation "is not completely tamper resistant and those intent on abusing this new formulation will likely find a means to do so. In addition, the product can still be misused or abused and result in overdose by simply administering or ingesting larger than recommended oral doses."⁷³⁰

1042. Despite this knowledge, the Opioid Marketing Enterprise pursued messaging and a strategy that was deceptive and was designed to deceive doctors in particular. Even after Purdue pleaded guilty to offenses related to its marketing and distribution of addictive opioids, Defendants and McKinsey advised Purdue to market OxyContin to encourage more prescriptions (that it knew would lead to abuse and overdose events) into higher dose prescriptions by a smaller number of loyalist prescribers.

1043. Rather than the deception of doctors being an unforeseen consequence, Defendants and McKinsey intentionally set out to target doctors as a cog in the Enterprise's scheme to defraud. Indeed, deceiving doctors was part of the marketing scheme, and doctors were utilized in furtherance of the marketing scheme. Medical providers were not a break in the causal chain of

⁷³⁰ FDA Summary Review, https://www.accessdata.fda.gov/drugsatfda_docs/nda/2010/022272s000SumR.pdf

harm to Plaintiffs but were targeted players in the scheme to defraud and key links in the casual chain.

1044. The marketing scheme involved using data to target high prescribers and training marketers to make misleading statements with the goal to increase high dose prescriptions which Defendants and McKinsey and Opioid Marketing Enterprise Members knew were more likely to be abused. Enterprise Members knew that overdoses were expected and that such overdoses would lead to need for increased services.

1045. Purdue's 2020 guilty plea acknowledged its role in using aggressive marketing to convince doctors to prescribe opioids unnecessarily, fueling the drug addiction crisis. Defendants and McKinsey were the masterminds of marketing scheme following Purdue's 2007 guilty plea. Defendants and McKinsey developed and helped implement these strategies.

1046. In an October 26, 2009 presentation, "OxyContin – driving growth through stronger brand loyalty," McKinsey proposed tactics to turnaround declining sales, "[e]nhance loyalty to OxyContin among loyalist prescribers," "convert[ing] 'fence sitters' into more loyal OxyContin prescribers," and "protect OxyContin's market share[.]" In other words, McKinsey proposed increasing sales by pushing both willing and reluctant physicians to prescribe more OxyContin. Defendants worked assiduously alongside McKinsey to accomplish these goals.

1047. Defendants and McKinsey recommended segmenting prescribers and tailoring messages and tactics to different segments. For prescribers dubbed "Early Adopting Experts" and "Proactive Teachers," defined by a willingness to use extended release opioids, including in patients who were not already using opioids, McKinsey urged emphasizing that its 7 tablet strengths provide flexibility to "tailor the dose" to customer needs.⁷³³ Upon information and belief, this

⁷³¹ MCK-MDL2996-0126522

⁷³² *Id.* at 2

⁷³³ *Id.* at 12.

1048. Purdue adopted McKinsey's

message aimed to encourage prescribers to initiate and maintain patients on OxyContin long-term by reminding them they could increase the dose as patients became tolerant with long-term use (rather than discontinue use when the drug lost its effectiveness).

Defendants Publicis and ZS were integral to these efforts, and Publicis worked for years to continually refine titration messaging for Purdue to ensure each prescription was as profitable as possible.

1049. As detailed throughout, Defendants and McKinsey and Opioid Marketing Enterprise Members were aware of the catastrophic injury inflected on the public by selling harmful, addictive opioid products. Yet when promoting opioids and engaging in doctor detailing, the Enterprise Members intentionally hid the potential for abuse and addiction by marketing OxyContin's 12-hour dosing as meaning that users only need to take OxyContin twice a day, thus requiring fewer pills.

1050. It was foreseeable that this marketing strategy would lead to greater addiction because OxyContin wore off after 8 to 10 hours in many patients. Prescribing 12-hour dosing led to "end of dose failure," which led to a vicious cycle that became "the perfect recipe for addiction." As a result, what Defendants and McKinsey marketed as "convenient" led to what was described as "a [d]escription of Hell."

⁷³⁴ PPLPC023000251226 (see also PPLPC012000243668 (PPLPC012000245087 PPLPC012000246009 (PPLPC021000265092 PPLPC021000265092

⁷³⁵ PKY183123435

⁷³⁶ Harriet Ryan, ""You Want a Description of Hell?" OxyContin's 12-Hour Problem," Los Angeles Times, May 5, 2016, available at http://www.latimes.com/projects/oxycontin-part1/.
⁷³⁷ Id

1051. The marketing scheme worked. Nationwide, based on an analysis by the Los Angeles Times, more than 52% of patients taking OxyContin longer than three months are on doses greater than 60 milligrams per day—which converts to the 90 morphine equivalent dose that the CDC Guideline urges prescribers to "avoid" or "carefully justify."

1052. A key element of the marketing scheme that fueled the deadly epidemic of opioid abuse was doctor detailing using detailed prescriber data.

<u>Data Scheme: Use of Prescriber Data for Intentional Targeting of High Opioid Prescribers</u> Not Diversion Prevention

1053. Defendants and McKinsey were advisors to DEA registrants and Opioid Marketing Enterprise Members, who had a legal duty to guard against diversion and report suspicious orders of controlled substances. Rather than assisting in reporting suspicious orders, Defendants and McKinsey used their position and access to detailed prescriber information to actually divert resources to target high volume prescribers to sell more opioids.

1054. Distributors of controlled substances have a legal duty to report suspicious orders, and to report those that deviate substantially from a normal pattern and orders of unusual size and frequency. See 21 U.S.C. § 823; 21 C.F.R. § 1301.74(b). These obligations included a legal duty to maintain effective controls and procedures to guard against diversion of controlled substances and a legal duty to maintain a system to identify and report suspicious orders of controlled substances. See 21 C.F.R. §§ 1301.7(a) (b); 1301.74(b). Rather than advising their registrant clients on how to comply with their legal duties to maintain effective controls to guard against diversion and how to operate a system to identify and report suspicious orders, in furtherance of the scheme, Defendants and McKinsey and the Opioid Marketing Enterprise Members used detailed data to target prescribers to increase the opioid market.

⁷³⁸ CDC Guideline at 16.

1055. Consistent with the Enterprise's purpose of increasing profit by deceptively marketing opioids, McKinsey was tasked with "Identifying Granular Growth Opportunities for OxyContin," conducting an "assessment of the underlying drivers of current OxyContin performance," identifying "key opportunities to drive near-term OxyContin performance," and developing "plans to capture priority opportunities." Defendants assisted and collaborated with McKinsey to identify and exploit these growth opportunities and drivers of near-term performance, and implemented plans to capture these opportunities.

1056. Defendants and McKinsey received physician-level sales data to develop its marketing strategy to increase OxyContin performance after Purdue's 2007 guilty plea. Rather than using this access to the granular data to avoid diversion and to prevent Enterprise members from targeting prescribers with suspicious prescribing patterns, Defendants and McKinsey used this information to help the Opioid Marketing Enterprise members push more opioids on high volume prescribers in furtherance of its schemes to defraud. The targets were chosen based on their history of prescribing high doses of opioids in large quantities.

1057. One of the services the Enterprise used in furtherance of this scheme concerned the use of data to help Purdue meet its goals. Defendants and McKinsey's analysis for "Evolve to Excellence" shows that Defendants and McKinsey had detailed information from which it could discern, as could Purdue, whether a prescriber had problematic patterns suggesting operation as a "pill mill," including a shift to other opioids after OxyContin's reformulation. Yet, Defendants and McKinsey urged Purdue to target, and seek to increase the prescribing of, all of these prescribers from whom it perceived Purdue could obtain greater profits.

1058. McKinsey found that Purdue did not "focus on the highest potential docs," measured both by the number of prescriptions and reimbursement considerations. A McKinsey analyst

⁷³⁹ PPLPC030000770531

⁷⁴⁰ MCK-MDL2996-0364024

urged McKinsey to recommend Purdue target "[1]iterally, at least all" prescribers in the top 20% of prescribers, "minus another few percent who are no sees[.]" McKinsey team lead Arnab Ghatak replied that "they probably have 20% no see[], but i'd also assume there are not many high writers that are no see." ("No see" prescribers are prescribers who do not accept visits from pharmaceutical sales representatives. Thus, upon information and belief, McKinsey recognized that most of the highest volume prescribers, or "high writers" of prescriptions, were willing to entertain sales visits from sales representatives.) Defendant ZS assisted McKinsey with targeting these prescribers, and Publicis assisted McKinsey with crafting messages to deliver to them.

1059. The Opioid Marketing Enterprise used data for intentional targeting of high prescribers and not for diversion prevention. Defendants and McKinsey advised Purdue to raise sales of Oxycontin by focusing on high dose sales and deceptively messaging to physicians that OxyContin would improve function and quality of life. Defendants and McKinsey urged Purdue to maximize sales by dictating which prescribers its sales representatives would target. For example, McKinsey advised Purdue that it should take "specific actions" to increase sales of OxyContin, including "Prescriber Targeting" and "Turbocharg[ing] Purdue's Sales Engine." Defendant ZS's work was particularly relevant to these efforts.

1060. Defendants and McKinsey targeted not just doctors but also nurse practitioners and physician assistants, with McKinsey recommending Purdue "[d]ouble down on nurse practitioners and physician assistants . . . as they represent a growing market segmentation of prescribers."⁷⁴²

1061. The Enterprise's scheme also explored ways to increase the amount of time sales representatives spent in the field increasing opioid sales, and prioritizing OxyContin in incentive compensation targets.⁷⁴³ Again, ZS's work was particularly relevant to these efforts.

⁷⁴¹ MCK-MDL2996-0364267

⁷⁴² MCK-MDI 2996-0303399

⁷⁴³ PPLPC012000437346

1062. By April 24, 2014, the plan was working and McKinsey reported that Purdue's "sales force is selecting an increasing percentage of high-value OxyContin prescribers as targets."

1063. McKinsey ensured Purdue would benefit from the lessons learned by other Enterprise members, stating that "its experience with other pharmaceutical companies suggests that such a comprehensive Sales transformation program takes nine months." Likewise, McKinsey recommended physician targeting to other Enterprise members, including Endo and Janssen. Similarly, ZS and Publicis were also advising multiple Opioid Marketing Enterprise members at the same time.

Enterprise was working toward the common purpose of deceptively convincing doctors to prescribe more opioids and thereby increase their own profits. By developing "Evolve to Excellence," which was implemented as a plan to "turbocharge" opioid sales, McKinsey advised that Purdue would see a greater return on its sales investment by focusing its targets, including on prescribers with alarming prescribing patterns that raised red flags they were writing "prescriptions" for non-medical use. Defendants worked for years with McKinsey to accomplish these goals. The plan aimed at boosting sales of OxyContin by targeting the highest volume opioid prescribers, which Defendants and McKinsey and the other members of the Opioid Marketing Enterprise knew and/or should have known would result in the expansion of the illicit opioid market.

1065. The Enterprise sought to grow opioid sales to prescribers who raised red flags of diversion and orders it knew or should have known were likely to be diverted or fuel an illegal market. Purdue had a legal obligation not to target these prescribers; rather, it was obligated to

⁷⁴⁴ MCK-MDL2996-0104840; PPLPC035000220406

⁷⁴⁵ MCK-MDI 2996-0187168

⁷⁴⁶ MCK-MDL2996-0130803; MCK-MDL2996-0135713

report their conduct to law enforcement. Yet the Enterprise used access to prescriber data not to report diversion but to enhance diversion.

Pattern of Racketeering Activity

1066. Defendants and McKinsey together with the other Opioid Marketing Enterprise Members engaged in a scheme to unlawfully increase sales of opioids—and grow their share of the prescription painkiller market—through repeated and systematic misrepresentations about the safety and efficacy of opioids for treating long-term chronic pain. As a unique consulting entity with knowledge of both the addictive properties and abuse potential of opioids and with access to data regarding internal prescribing behaviors of its targets, McKinsey perpetrated – with Defendants ongoing aid and assistance – a number of fraudulent schemes using the mails and wires, including advising Purdue to market more opioids, in higher doses, to high volume prescribers while helping Purdue avoid mandatory prescriber education regarding the risks of opioids. Defendants and McKinsey fueled the epidemic alongside their clients. Through targeted marketing that Defendants and McKinsey developed, "turbocharged," and implemented, Defendants and McKinsey substantially contributed to an explosion in the use of opioids across the United States. Defendants and McKinsey are engaged in and affect interstate commerce because they advised multiple opioid manufacturers headquartered on different states on the sale of opioid products across the United States, as alleged herein.

1067. The Opioid Marketing Enterprise Members devised and knowingly carried out this illegal scheme and artifice to defraud by means of materially false or fraudulent pretenses, representations, promises, or omissions of material facts regarding the safe, non-addictive and effective use of opioids for long-term chronic, non-acute, and non-cancer pain. They knew that these representations deviated from the FDA-approved use of these drugs and were not supported by actual evidence. The Opioid Marketing Enterprise Members intended that their common purpose

and scheme to defraud would, and did, deceive consumers, prescribers, regulators, Plaintiffs, and other intended victims and they used the U.S. Mail and interstate wire facilities with the specific intent to advance, and for the purpose of executing, their illegal scheme.

1068. By intentionally concealing the material risks and affirmatively misrepresenting the benefits of using opioids for chronic pain, the Opioid Marketing Enterprise Members engaged in a fraudulent and unlawful course of conduct constituting a pattern of racketeering activity.

1069. To achieve the common goal and purpose of the Opioid Marketing Enterprise, the Opioid Marketing Enterprise Members hid from the consumers, prescribers, regulators, and Plaintiffs: (a) the fraudulent nature of the Opioid Marketing Enterprise Members' marketing scheme; (b) the fraudulent nature of statements made by the Opioid Marketing Enterprise Members regarding the safety and efficacy of prescription opioids; and (c) the true nature of the relationship between the members of the Opioid Marketing Enterprise.

1070. The Opioid Marketing Enterprise Members with knowledge and intent, to the overall objective of the Opioid Marketing Enterprise Members' fraudulent scheme and participated in the common course of conduct to commit acts of fraud and indecency in marketing prescription opioids.

1071. Indeed, for the Opioid Marketing Enterprise Members' fraudulent scheme to work, each of them had to agree to implement similar tactics regarding fraudulent marketing of prescription opioids. This coordination was accomplished via their relationships with each other and via Defendants and McKinsey's relationships and contacts with key opioids manufacturers.

1072. The Opioid Marketing Enterprise Members' predicate acts all had the purpose of creating the opioid epidemic that substantially injured Plaintiffs, while simultaneously generating billion-dollar revenues and profits for the Opioid Marketing Enterprise Members. The predicate acts were committed or caused to be committed by the Opioid Marketing Enterprise Members

through their participation in the Opioid Marketing Enterprise and in furtherance of its fraudulent scheme.

1073. The Opioid Marketing Enterprise Members' scheme described herein was perpetrated, in part, through multiple acts of mail fraud and wire fraud, constituting a pattern of racketeering activity. Defendants and McKinsey used mail and wire transmission, directly or indirectly, in furtherance of this scheme by transmitting deliberately false and misleading statements to prescribers and the public.

1074. Defendants and McKinsey had a specific intent to deceive and defraud prescribers, regulators and Plaintiffs. For example, as alleged above, Defendants and McKinsey made repeated and unequivocal statements through the mails and wires that were false and misleading. For example, McKinsey advised Purdue to market OxyContin based on the false and misleading notion that the drug can provide "freedom" and "peace of mind" for its users, and concomitantly reduce stress and isolation.

1075. Similarly, they caused to be transmitted through the mails and wires false and misleading statements regarding the addiction potential of opioids. Moreover, Defendants and McKinsey had direct involvement in marketing statements and thus caused the statements to be made, notwithstanding that they knew they were false for the reasons detailed above.

1076. The marketing scheme is especially egregious since the public relies on physicians as a position of trust and authority in the community regarding their health and well-being. Defendants and McKinsey intentionally deceived physicians regarding the abuse potential of opioids. It intended prescribers and the public to rely on its false statements. Defendants and McKinsey intended reliance on these false statements as it was their goal for doctors to prescribe more and higher quantities of these dangerous pills to the public. This scheme was therefore

reasonably calculated to deceive not only persons of ordinary prudence and comprehension but also educated physicians in a place of high trust in the community.

Predicate Acts

- 1077. To carry out, or attempt to carry out, the scheme, the Enterprise Members, each of whom is a person associated-in-fact with the Enterprise, did knowingly conduct or participate in, directly or indirectly, the affairs of the Enterprise through a pattern of racketeering activity within the meaning of 18 U.S.C. §§ 1961(1), 1961(5) and 1962(c), and employed the use of the mail and wire facilities, in violation of 18 U.S.C. § 1341 (mail fraud) and § 1343 (wire fraud).
- 1078. Specifically, the Enterprise Members have committed, conspired to commit, and/or aided and abetted in the commission of, at least two predicate acts of racketeering activity (i.e., violations of 18 U.S.C. §§ 1341 and 1343), within the past ten years.
- 1079. The multiple acts of racketeering activity which the Enterprises Members committed, or aided or abetted in the commission of, were related to each other, posed a threat of continued racketeering activity, and therefore constitute a "pattern of racketeering activity."
- 1080. The racketeering activity was made possible by the Enterprise's regular use of the facilities, services, distribution channels, and employees of the Enterprise Members.
- 1081. The Opioid Marketing Enterprise Members participated in the schemes by using mail, telephone, and the internet to transmit mailings and wires in interstate or foreign commerce.
- 1082. The Enterprise Members used, directed the use of, and/or caused to be used, thousands of interstate mail and wire communications in service of their schemes through common misrepresentations, concealments, and material omissions.
- 1083. In devising and executing the illegal schemes, the Opioid Marketing Enterprises Members devised and knowingly carried out a material scheme and/or artifice to defraud Plaintiffs

and prescribers and to obtain money by means of materially false or fraudulent pretenses, representations, promises, or omissions of material facts.

1084. For the purpose of executing the illegal schemes, the Enterprise Members committed these racketeering acts, which number in the thousands, intentionally and knowingly with the specific intent to advance the illegal schemes.

1085. The Opioid Marketing Enterprise Members' predicate acts of racketeering (18 U.S.C. § 1961(1)) include, but are not limited to the conduct described in the Factual Allegations section of this Complaint, and:

1086. Mail Fraud: The Opioid Marketing Enterprise Members violated 18 U.S.C. § 1341 by sending or receiving, or by causing to be sent and/or received, materials via U.S. mail or commercial interstate carriers for the purpose of executing the unlawful scheme to design, manufacture, market, and sell the prescription opioids by means of false pretenses, misrepresentations, promises, and omissions.

1087. Wire Fraud: The Opioid Marketing Enterprise Members violated 18 U.S.C. § 1343 by transmitting and/or receiving, or by causing to be transmitted and/or received, materials by wire for the purpose of executing the unlawful scheme to design, manufacture, market, and sell the prescription opioids by means of false pretenses, misrepresentations, promises, and omissions.

1088. The Opioid Marketing Enterprise Members' use of the U.S. Mail and interstate wire facilities to perpetrate the opioids marketing scheme involved thousands separate instances of the use of the U.S. Mail or interstate wire facilities in furtherance of the unlawful Opioid Marketing Enterprise, including essentially uniform misrepresentations, concealments, and material omissions regarding the beneficial uses and non-addictive qualities for the long-term treatment of chronic, non-acute, and non-cancer pain, with the goal of profiting from the increased sales of the Opioid Marketing Enterprise Members' drugs that occurred because consumers, prescribers, regulators,

and Plaintiffs relied on the Opioid Marketing Enterprise Members' misrepresentations. These uses of the U.S. Mail or interstate wires included, inter alia:

- 1089. Marketing materials about opioids and their risks and benefits, which the Opioid Marketing Enterprise Members sent to health care providers, transmitted through the internet and television, and published across the country, including in counties and cities and to Plaintiffs;
- 1090. Written representations and telephone calls among the Opioid Marketing Enterprise Members and between the Opioid Marketing Enterprise Members regarding the misrepresentations, marketing statements, and claims about opioids, including the non-addictive, safe use of opioids for chronic, long-term pain generally;
- 1091. E-mails, telephone calls, and written communications among the Opioid Marketing Enterprise Members agreeing to or implementing the opioids marketing scheme;
- 1092. Communications among the Opioid Marketing Enterprise Members and between the Opioid Marketing Enterprise Members and the media regarding the publication, drafting, and dissemination of treatment guidelines as part of the Opioid Marketing Enterprise;
- 1093. Written and oral communications directed to prescribers, the public, and Plaintiffs that fraudulently misrepresented the risks and benefits of using opioids for chronic pain; and
- 1094. Receipts of increased profits sent through the U.S. Mail and interstate wire facilities—the wrongful proceeds of the scheme.
- 1095. Many of the precise dates of the fraudulent uses of the U.S. mail and interstate wire facilities are not obtainable (e.g., each time a McKinsey trained marketer "calls" or reached out to a physician targeted by ZS with materials designed and prepared by Publicis using the mails or wires in furtherance of the marketing scheme). Because the Opioid Marketing Enterprise Members disguised their participation in the Enterprise, and worked to keep the Enterprise's existence secret, many of the precise dates of the Opioid Marketing Enterprise's uses of the U.S. Mail and interstate

wire facilities (and corresponding predicate acts of mail and wire fraud) have been hidden and cannot be alleged without access to the books and records maintained by the Opioid Marketing Enterprise Members. Indeed, an essential part of the successful operation of the Opioid Marketing Enterprise alleged herein depended upon secrecy. Plaintiffs have, however, described the types of predicate acts of mail and/or wire fraud, including the specific types of fraudulent statements upon which, through the mail and wires, Defendants and McKinsey engaged in fraudulent activity in furtherance of their scheme.

1096. The factual allegations in this Complaint describes multiple occasions of Defendants working to create and deliver to targeted audiences, including no-see prescribers, misrepresentations and false statements in furtherance of the scheme. Below, Plaintiffs also describe examples of occasions on which other Opioid Marketing Enterprise Members disseminated misrepresentations and false statements to consumers, prescribers, regulators, and Plaintiffs, and how those acts were also in furtherance of the scheme.

From	To	Date	Description
Purdue	Prescribers and Plaintiffs	2007	Statements that pain relief from opioids improves patients' function and quality of life in advertising and a book
Purdue	Prescribers	Continuous	Telephonic and electronic communications by its sales representatives indicating that opioids will improve patients' function
Purdue	FDA advisory committee	September 2009	Presentation prepared by McKinsey indicating that its reformulated OxyContin will deter abuse
Purdue	Prescribers and Plaintiffs	2010 onwards	Statements that the reformulated OxyContin will deter abuse and therefore doctors can continue to safely prescribe opioids
Purdue	Prescribers and Plaintiffs	2010-2020	Statements from Purdue at McKinsey's direction that opioids can provide "freedom," "peace of mind," and give patients "the best possible chance to live a full and active life"
Purdue	Prescribers and Plaintiffs	Advertising produced in 2016	Advertising from Purdue that "We sell hope in a bottle."
Purdue	Prescribers and Plaintiffs	2010 onwards	Statements that OxyContin's 12-hour dosing would allow patients to only need to take OxyContin twice a day, thus requiring fewer pills

1	Purdue	Prescribers	2013	Statements from Purdue at McKinsey's direction that
		and	onwards	OxyContin allowed physicians to "Individualize the Dose"
2		Plaintiffs		and that the dose of OxyContin can safely be increased or
3	T 1	D '1	2000	tailored as the patients adapt to a certain dose
	Endo	Prescribers and	2009	Statements made on an Endo-sponsored website,
4		Plaintiffs		PainKnowledge.com, indicating that patients who take opioids as prescribed usually do not become addicted
_	Endo	Prescribers	2009	Statements made on another Endo-sponsored website,
5	Lindo	and	2007	PainAction.com, indicating that most chronic pain patients do
6		Plaintiffs		not become addicted to opioid medications
0	Endo	Prescribers	Various	Statements in pamphlets and publications described by Endo
7		and		indicating that most people who take opioids for pain relief do
		Plaintiffs		not develop an addiction
8	Endo	Prescribers	Various	Statements made on the Endo-run website, Opana.com,
		and		indicating that opioid use does not result in addiction
9	Endo	Plaintiffs Prescribers	Various	Statements made on the Ende min website. Once com
10	Elido	and	various	Statements made on the Endo-run website, Opana.com, indicating that opioid dependence can be addressed by dosing
		Plaintiffs		methods such as tapering
11	Endo	Prescribers	Various	Statements made on its website, PainKnowledge.com, that
12		and		opioid dosages could be increased indefinitely
12		Plaintiffs		, , ,
13	Endo	Prescribers	Various	Statements made in a publication entitled "Understanding
		and		Your Pain: Taking Oral Opioid Analgesics" suggesting that
14	77.1	Plaintiffs	** .	opioid doses can be increased indefinitely
	Endo	Prescribers	Various	Electronic and telephonic communications to its sales
15				representatives indicating that the formula for its medicines is "crush resistant"
16	Endo	Prescribers	2007	Statements that pain relief from opioids improves patients'
		and		function and quality of life in advertising and a book
17		Plaintiffs		1 7
10	Endo	Prescribers	Various	Telephonic and electronic communications by its sales
18				representatives indicating that opioids will improve patients'
19	<u> </u>		** .	function
17	Janssen	Prescribers	Various	Statements on its website, PrescribeResponsibly.com,
20		and Plaintiffs		indicating that concerns about opioid addiction are overestimated
21	Janssen	Prescribers	2009	Statements in a 2009 patient education guide claiming that
21	b dires or	and	200)	opioids are rarely addictive when used properly
22		Plaintiffs		
	Janssen	Prescribers	2009	Statements included on a 2009 Janssen-sponsored website
23		and		promoting the concept of opioid pseudoaddiction
_		Plaintiffs		
24	Janssen	Prescribers	Various	Statements on its website, PrescribeResponsibly.com,
25		and		advocating the concept of opioid pseudoaddiction
23	Janssen	Plaintiffs Prescribers	Various	Statements on its website, PrescribeResponsibly.com,
26	Janssen	and	v arious	indicating that opioid addiction can be managed
		Plaintiffs		more and optota addiction can be managed
27	Janssen	Prescribers	2009	Statements in its patient education guide indicating the risks
28		and		associated with limiting the dosages of pain medicines
۵۵		Plaintiffs		

July 18,

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McKinsey

Purdue

offices to fraudulently promote OxyContin, including via (with 2013 "phone, video and even Google like proprietary tools" 747 prescribers as the planned target) April $2\overline{4}$, McKinsey Purdue Plan to promote OxyContin to "no-see" physicians through "remote interactions" including presenting "brand interaction and materials" "over the phone/internet" "748 (with 2017 prescribers as the planned target) McKinsey McKinsey July 14, Internal emails interpreting "the Purdue situation" and 2013 discussing OxyContin sales strategy including sales benchmarks and "focus on the highest potential docs" 749 Evolve 2 Excellence PowerPoint planning execution of the McKinsey Purdue September 23, 2013 (with scheme and discussing targeted performance metrics including "sales management calls per day, calls per year and prescribers adhering to target list",750 as the planned target) McKinsey July 30, Presentation showing "Scope of potential OxyContin growth Purdue opportunities" with proposed process including "Generate 2013 target list" and using "Reps/DMs [to] perform call planning (including refining target list)"⁷⁵¹

Discussion of McKinsey plan to increase calls to doctors'

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1097. Each of these fraudulent mailings and interstate wire transmissions constitutes racketeering activity and collectively, these violations constitute a pattern of racketeering activity, through which the Opioid Marketing Enterprise Members defrauded and intended to defraud consumers, prescribers, regulators, Plaintiffs, and other intended victims.

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1098. These were not isolated incidents. Instead, the Opioid Marketing Enterprise Members engaged in a pattern of racketeering activity by committing thousands of predicate acts in a five-year period, in the form of mail and wire fraud, and there remains a threat that such conduct will continue in the future.

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⁷⁴⁷ MCK-MDL2996-0104431, at 0104442

^{27 | &}lt;sup>748</sup> MCK-MDL2996-0104840

⁷⁴⁹ MCK-MDL2996-0364024

⁷⁵⁰ MCK-MDL2996-0316833, at 0316834

⁷⁵¹ MCK-MDL2996-0303399

1099. Each instance of racketeering activity alleged herein was related, had similar purposes, involved the same or similar participants and methods of commission, and had similar results affecting similar victims, including consumers, prescribers, regulators, and Plaintiffs. The Opioid Marketing Enterprise Members calculated and intentionally crafted the scheme and common purpose of the Opioid Marketing Enterprise to ensure their own profits remained high. In designing and implementing the scheme, the Opioid Marketing Enterprise Members understood and intended that those in the opioid distribution chain rely on the integrity of the pharmaceutical companies and ostensibly neutral third parties to provide objective and scientific evidence regarding the Opioid Marketing Enterprise Members' products.

1100. Opioid Marketing Enterprise Members' pattern of racketeering activity alleged herein and the Opioid Marketing Enterprise are separate and distinct from each other. Likewise, the Opioid Marketing Enterprise Members are distinct from the Opioid Marketing Enterprise.

amounted to a common course of conduct, with a similar pattern and purpose, intended to deceive consumers, prescribers, regulators, and Plaintiffs. Each separate use of the U.S. Mail and/or interstate wire facilities employed by the Opioid Marketing Enterprise was related, had similar intended purposes, involved similar participants and methods of execution, and had the same results affecting the same victims, including consumers, prescribers, regulators, and Plaintiffs. The Opioid Marketing Enterprise Members have engaged in the pattern of racketeering activity for the purpose of conducting the ongoing business affairs of the Opioid Marketing Enterprise.

1102. Each of the Opioid Marketing Enterprise Members aided and abetted others in the violations of the above laws, thereby rendering them indictable as principals in the 18 U.S.C. §§ 1341 and 1343 offenses.

1103. As described herein, the Opioid Marketing Enterprise Members engaged in a pattern of related and continuous predicate acts for many years. The predicate acts constituted a variety of unlawful activities, each conducted with the common purpose of obtaining significant money and revenue from the marketing and sale of their highly addictive and dangerous drugs. The predicate acts also had the same or similar results, participants, victims, and methods of commission. The predicate acts were related and not isolated events.

1104. The Opioid Marketing Enterprise Members' violations of law and pattern of racketeering activity directly and proximately caused Plaintiffs injury in their business and property. The Opioid Marketing Enterprise Members' pattern of racketeering activity logically, substantially, and foreseeably caused an opioid epidemic. The injuries of Plaintiff, as described herein, were not unexpected, unforeseen, or independent. Rather, as Plaintiffs allege, the Opioid Marketing Enterprise Members as a whole and, and Defendants and McKinsey in particular, knew that the opioids were unsuited to treatment of long-term chronic, non-acute, and non-cancer pain, or for any other use not approved by the FDA, and knew that opioids were highly addictive and subject to abuse. Nevertheless, the Opioid Marketing Enterprise Members engaged in a scheme of deception that utilized the mail and wires in order to carry out the Opioid Marketing Enterprise's fraudulent scheme, thereby increasing sales of their opioid products.

1105. It was foreseeable and expected that the Opioid Marketing Enterprise Members creating and then participating in the Opioid Marketing Enterprise through a pattern of racketeering activities to carry out their fraudulent scheme would lead to a nationwide opioid epidemic, including increased opioid addiction and overdose and the injuries that occurred as a result.

The Enterprise Was Well Aware of Risks of Abuse Before It "Turbocharged" its Marketing Scheme.

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1106. These devastating results were eminently foreseeable by the Opioid Marketing Enterprise Members.

1107. When Purdue pleaded guilty in 2007, it was evident that Purdue's behavior and excessive prescribing was directly linked to a drug addiction crisis that caused severe and extensive damage to America. Purdue's methods included "using aggressive marketing tactics to convince doctors to unnecessarily prescribe opioids – frivolous prescriptions that experts say helped fuel a drug addiction crisis that has ravaged America for decades."⁷⁵²

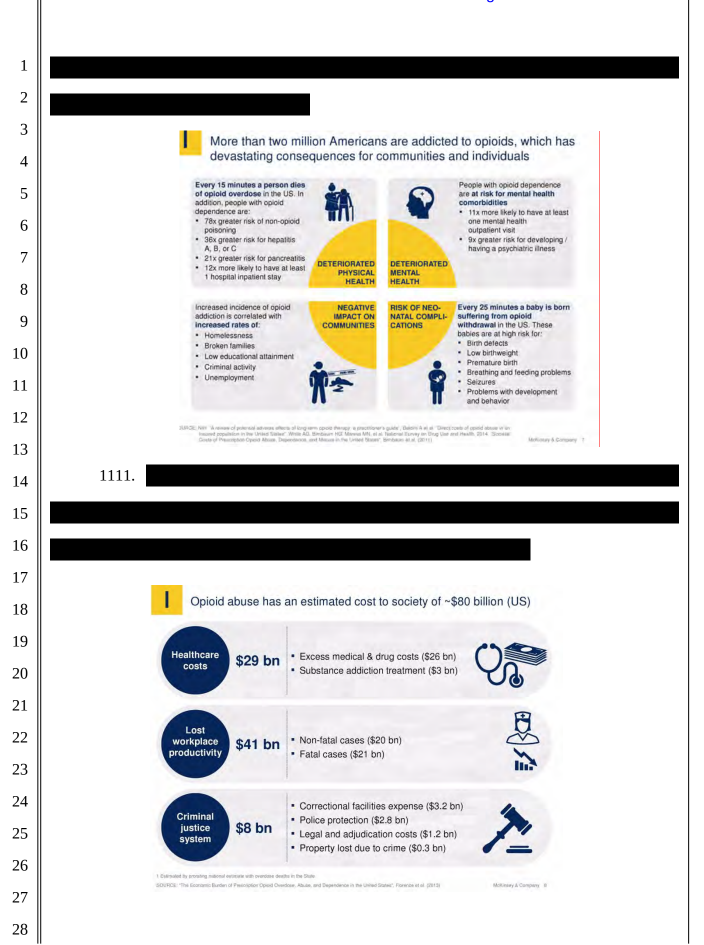
1108. Defendants and McKinsey cannot deny knowledge regarding Purdue's 2007 guilty plea. At that point, McKinsey knew that opioids were addictive. McKinsey knew that OxyContin was being widely abused and causing harm to people and entities like Plaintiffs. And McKinsey knew that Purdue had been fraudulently marketing OxyContin as less addictive, less subject to abuse, and less likely to cause withdrawal. And yet, years later, in 2013, McKinsey orchestrated a scheme with Defendants to continue to aggressively promote opioids despite knowledge that people were still dying from overdoses.

1109. Thus, Defendants and McKinsey continued to add fuel to this fire by persisting in aggressively marketing to physicians and continuing to fuel the opioid crisis after Purdue's guilty plea. It was foreseeable that continuing to do so would devastate American communities.



⁷⁵² Jan Hoffman & Katie Benner, *Purdue Pharma Pleads Guilty to Criminal Charges for Opioid Sales*, N.Y. Times (updated Dec. 17, 2020), https://www.nytimes.com/2020/10/21/health/purdue-opioids-criminal-charges.html.

⁷⁵³ MCK-MDL2996-0070516, at 0070517.



1112. Similarly, news stories across the nation reported additional consequences of wide scale opioid addiction: needles littered around public property, posing costs to the governments and danger to residents.⁷⁵⁴

1113. The foreseeability of the abuse and need for additional services that would be required following the misleading marketing and increased prescribing and use of high dose opioids is also evidenced by McKinsey's attempt to put a price tag on overdoses. McKinsey suggested payment amounts for event-based contracts: \$6,000 to \$15,000 (paid to health insurers for increased medical services). Indeed, McKinsey was well aware that increased prescriptions would lead to overdoses and to an additional financial burden for social and health services.

1114. Defendants and McKinsey are liable for their successful efforts to increase OxyContin sales after Purdue's 2007 guilty plea for misbranding the drug. Indeed, Defendants' and McKinsey's focus on increasing opioid sales after Purdue's guilty was incendiary to escalating and perpetuating the opioid epidemic by: (a) using data to specifically target high volume prescribers; (b) persuading sales of higher doses of opioids; (c) tailoring marketing messages to conceal their addictive principles; and (d) by reducing the training of sales representatives.

1115. In 2013, when the consent decree expired (which obligated Purdue to submit annual compliance reports regarding its marketing), Defendants and McKinsey helped Purdue reengage in its nefarious conduct of targeting and deceiving doctors about the abuse potential of opioids.

1116. After Purdue's guilty plea, Defendants and McKinsey identified physicians—that had already been influenced by Purdue's misrepresentations and were thus already high prescribers—as optimal targets for a massive marketing push to sell more OxyContin. Defendants and McKinsey monitored the prescription behaviors of individual doctors and utilized the

⁷⁵⁴ See, e.g., https://www.bostonglobe.com/metro/regionals/south/2014/10/25/hypodermic-needles-litter-landscape-south-boston/pzgmgbyjYFCD967TePDyiM/story.html

prescriber-level data and urged Purdue to allocate its time and resources to high prescribing physicians.

1117. By November 2013, Defendants and McKinsey had obtained the physician-level data it had previously requested and continued to study ways to sell additional OxyContin prescriptions by refining and targeting the sales pitch to them.

1118. In 2013, Project Turbocharge began. McKinsey proposed Project Turbocharge, a marketing strategy to increase opioids sales by hundreds of millions of dollars annually. With Defendants and McKinsey's ongoing aid and assistance, Purdue trained its sales representatives to operate using McKinsey's strategy for selling OxyContin. It is not coincidental to the Enterprise scheme that as soon as the constraints associated with its guilty plea and consent agreement ended, Defendants and McKinsey assisted Purdue in turbocharging sales.

1119. As Defendants and McKinsey were pushing hard to turbocharge and promote the sale of opioids, McKinsey anticipated and expected that people would die from opioid overdoses. It acknowledged this when, in 2017, it proposed that Purdue pay health insurers or other entities in the distribution chain rebates "for every OxyContin overdose attributable to pills they sold."

1120. Defendants and McKinsey cannot deny that it was not aware of the abuse and overdose potential of opioids when it provided estimates for the future costs of overdose or opioid use disorder events.

1121. Defendants and McKinsey and the other Opioid Marketing Enterprise Members marketed a product, through intentionally deceptive means, that it knew would result in consumer deaths and harm to Plaintiffs. This is not an attenuated causal chain. Rather, aggressively marketing to high prescribing individuals, and training to not fully disclose the risk of abuse, were integral

⁷⁵⁵ Walt Bogdanich & Michael Forsythe, *McKinsey Proposed Paying Pharmacy Companies Rebates for OxyContin Overdoses*, N.Y. Times (updated Nov. 5, 2021), https://www.nytimes.com/2020/11/27/business/mckinsey-purdue-oxycontin-opioids.html

parts of the marketing scheme. Publicis even described itself as the "strategic backbone" of McKinsey's Project Turbocharge initiative. Deceptive messaging to targeted prescribers who were likely to prescribe more pills in a dose with an anticipated abuse potential was part and parcel of the scheme to defraud.

- 1122. As a result, Plaintiffs have shouldered the burden of these anticipated increased services and harm to business and property that are inherently tied to opioid abuse and misuse, and both the increased services and harms were reasonably and actually expected from increased prescribing.
- 1123. The Enterprise's goal was to increase opioid prescribing, and the Enterprise Members knew that doing so would also result in the need for increased medical services. It was also foreseeable that increased prescriptions would also result in increased costs to Plaintiffs and communities throughout the United States.
- additional resources or suffered other harm to business and property as a result of harms associated with opioid addiction. The Enterprise persisted in targeting prescribers to prescribe high doses of opioids and knew that doing so would result in adverse health and social outcomes, including overdoses, neo-natal complications, harm to communities like Plaintiffs, hazardous waste in Plaintiff communities, as well as and increased expenditures on services to combat such ill effects.

Plaintiffs' Business and Property Have Been Damaged by the Enterprise's RICO Violations.

1125. The Opioid Marketing Enterprise's misleading marketing and failure to prevent prescription opioid diversion damaged Plaintiffs. In addition to medical services, the Opioid Marketing Enterprise's misconduct has contributed to a range of social problems, including violence and delinquency. Adverse social outcomes include child neglect, family dysfunction,

babies born addicted to opioids, criminal behavior, poverty, property damage, unemployment, and social despair. As a result, Plaintiffs are devoting more and more resources to the opioid epidemic.

- 1126. Notably, Plaintiffs have experienced vast harm to business and property directly, proximately, and foreseeably caused by the racketeering enterprise. The full extent of each Plaintiffs' damage cannot be fully captured in this pleading but can be fleshed out during the bellwether process. Below are some discrete examples that demonstrate the common and typical universal harm to Plaintiffs and the specific types of harm foreseeably caused by the Opioid Marketing Enterprise.
- 1127. Specifically, the Opioid Marketing Enterprise Members' creation of, and then participation in, the Opioid Marketing Enterprise through a pattern of racketeering activities to carry out their fraudulent scheme has injured Plaintiffs in the form of substantial losses of money and property that logically, directly and foreseeably arise from the opioid epidemic. The injuries to Plaintiffs, as alleged throughout this Complaint, and expressly incorporated herein by reference, include new, different, and increased expenditures by and losses to Plaintiffs, for example:
 - A. Hazardous waste in Plaintiffs' communities, including on Plaintiffs' real property;
 - B. Costs for providing healthcare and medical care, additional therapeutic, and prescription drug purchases, and other treatments for patients suffering from opioid-related addiction or disease, including overdoses and deaths;
 - C. Costs of training first responders in the proper treatment of drug overdoses;
 - D. Costs associated with providing first responders with naloxone—an opioid antagonist used to block the deadly effects of opioids in the context of overdose;
 - E. Costs associated with emergency responses by first responders to opioid overdoses;

F. Costs for providing mental health services, treatment, counseling, rehabilitation services, and social services to victims of the opioid epidemic and their families;

- G. Costs associated with the injuries to the health and welfare of the residents who reside in the jurisdiction of Plaintiffs caused by the opioid epidemic;
- H. Costs associated with providing care for children whose parents suffer from opioid-related disability or incapacitation; and
- I. Losses caused by the diversion of revenue to address the opioid epidemic that would otherwise have been used to provide other services.
- 1128. The injuries to Plaintiffs were directly and proximately caused by the Enterprise's racketeering activities because they were the logical, substantial, and foreseeable cause of the injuries to Plaintiffs. But for the opioid epidemic created by the Opioid Marketing Enterprise Members through their Opioid Marketing Enterprise, Plaintiffs would not have lost money or property, and the health and welfare of citizens would not have been harmed.
- 1129. Plaintiffs have been injured by the Enterprise's conduct, and such injury would not have occurred but for the predicate acts which also constitute acts taken in furtherance of the conspiracy pursuant to Section 1962(d). By working to expand the opioid market, fraudulently concealing the abuse potential of opioids, targeting high volume prescribers, and deceiving prescribers and the public in order to allow opioids to continue to remain on the market, the Enterprise caused the expansion of opioid prescribing and caused a large number of people across the United States, including in Plaintiffs' communities to become addicted to opioids, thus forcing Plaintiffs to expend, time, money and resources to address the opioid epidemic that Defendants and McKinsey and the Enterprise created through their conduct. Indeed, Defendants and McKinsey intentionally deceived doctors and public health workers in order to continue to grow the opioid

market. The repeated fraudulent misstatements by Defendants and McKinsey contributed to an explosion in the use of opioids across the country.

1130. Plaintiffs were direct victims of Defendants and McKinsey's misconduct. The Enterprise displayed a wanton disregard for public health and safety by intentionally deceiving doctors about the addiction potential of opioids and by marketing higher doses to physicians. The harm created by Defendants and McKinsey required Plaintiffs to expend financial and other resources to mitigate the health crisis of opioid misuse and addiction. The expansion of this market was the goal of the Enterprise and was critical to its success. Therefore, the harm suffered by Plaintiffs to their property forced them to expend resources beyond the ordinary costs of services to combat the opioid epidemic, was directly foreseeable, and in fact, was an intentional result of Defendants' and McKinsey's misconduct. Indeed, McKinsey anticipated overdose events and actually estimated price premiums on these expected overdose events. Defendants and McKinsey knew that the products it was marketing were highly addictive and could lead to deadly overdoses yet continued to "turbocharge" sales by fraudulently pushing the product on doctors through its deceptive marketing scheme.

1131. The creation and implementation of the marketing scheme that Defendants and McKinsey developed and deployed through its Enterprise directly harmed Plaintiffs by imposing costs on their businesses and properties. The harm caused by this scheme was an unnatural, human-caused, profit-driven, and completely preventable disaster (had the Enterprise Members obeyed the law). Thus, Plaintiffs' injuries are not solely the result of routine government expenses. Instead, as a result of Defendants' and McKinsey's misconduct, Plaintiffs have been and will be forced to go far beyond what a governmental entity might ordinarily be expected to pay to enforce laws and to promote the general welfare in order to combat the opioid epidemic, whose primary origins were in prescription opioids administered by prescribers whom Defendants and McKinsey targeted with

their marketing scheme to increase sales. This includes providing new programs and new services as a direct result and in direct response to Defendants' and McKinsey's misconduct. In addition, Plaintiffs have suffered losses to their property as a direct result of the kind of inevitable consequences of the drug addiction and criminal behavior that Defendants and McKinsey predicted. As a result of the conduct of the Enterprise, Plaintiffs have incurred and will continue to incur costs that far exceed the norm.

- 1132. The injuries to Plaintiffs were directly and proximately caused by these racketeering activities because they were the logical, substantial, and foreseeable cause of the injuries to Plaintiffs. But for the opioid epidemic the Opioid Marketing Enterprise Members created through their Opioid Marketing Enterprise, Plaintiffs would not have lost money or property, and the health and welfare of residents in Plaintiffs' jurisdictions would not have been harmed. Moreover, Defendants and McKinsey's internal documents show that it actually did foresee many of the harms that resulted from its conduct.
- Defendants' mail and wire fraud and Plaintiffs' injuries. Defendants, in furtherance of the Enterprise's common purpose, caused to be made false and misleading statements directly to the doctors (who consumers rely on to provide health advice) and the public. Doctors are not a break in the causal chain. Instead, the Enterprise members as a whole, and Defendants and McKinsey in particular, intentionally targeted doctors and sought to deceive them. That doctors were then deceived and behaved as the Enterprise wanted, prescribing more and more opioids, was the purpose of the scheme, not an intervening cause.
- 1134. The Enterprise's violations of 18 U.S.C. § 1962(c) have directly and proximately caused injuries and damages to Plaintiffs, and Plaintiffs are entitled to bring this action for three

times their actual damages, as well as for injunctive/equitable relief, costs and reasonable attorneys' fees and costs pursuant to 18 U.S.C. § 1964(c).

COUNT II: Negligence

- 1135. Plaintiff realleges and incorporates by reference the allegations set forth above.
- 1136. Negligence is established where the defendant owes the plaintiff a duty of care, breaches that duty, and the plaintiff sustained an injury or loss proximately caused by the defendant's breach.
- 1137. Defendants, through their work with Purdue and other opioid manufacturers, owed a duty of care to the Plaintiff, pursuant to which it would not encourage the over-marketing and over-prescribing of a controlled substance known at the time to be addictive and known at the time to be a threat to public health.
- 1138. In violation of this duty, for decades Defendants devised and assisted Purdue, and other opioid manufacturers, with implementing sales and marketing campaigns, including prescriber targeting and salesforce incentive compensation structures that would dramatically increase the amount of opioids prescribed and distributed in the Sac and Fox Nation.
- 1139. As a direct and proximate result of Defendants' negligent conduct, the Sac and Fox Nation has suffered and will continue to suffer harm.

COUNT III: Gross Negligence

- 1140. Plaintiff realleges and incorporates by reference the allegations set forth above.
- 1141. The oversupply of opioids and plague of addiction led to a widespread epidemic of overdoses, illness, and death that claimed thousands of lives and cost many millions of dollars of public spending—circumstances that constituted an imminent or clear and present danger amounting to more than normal and usual peril.

a duty of care to the Plaintiff, pursuant to which they would not encourage the over-marketing and over-prescribing of a controlled substance known at the time to be addictive and known at the time to be a threat to public health.

1143. In violation of this duty, for decades Defendants devised and assisted Purdue and other opioid manufacturers with implementing sales and marketing campaigns, including Purdue's *Evolve to Excellence* campaign, that would dramatically increase the amount of opioids prescribed and distributed to Plaintiff's citizens.

1144. As a direct and proximate result of Defendants' negligent conduct, Plaintiff has suffered and will continue to suffer harm.

COUNT IV: Negligent Misrepresentation

- 1145. Plaintiff realleges and incorporates by reference the allegations set forth above.
- 1146. Defendants, in the course of their business with Purdue and other opioid manufacturers, failed to exercise reasonable care or competence by communicating false information regarding Defendants' clients' opioids that Defendants knew would be used for guidance by others in their business transactions, including the healthcare providers within Plaintiff who were capable of prescribing Purdue's drugs.
- 1147. Plaintiff is one of a limited group of entities to whom Defendants knew Purdue and other opioid manufacturers intended to supply with false information regarding opioids.
- 1148. Defendants knew that the false information was material to healthcare providers' decision to prescribe opioids to patients. Defendants intended that such statements be relied upon to encourage additional opioid prescriptions.

COUNT V: Public Nuisance

1149. Plaintiff realleges and incorporates by reference the allegations set forth above.

1150. Oklahoma courts have long recognized that anything which interferes with public health and promotes the spread of disease, bodily injury, and/or death, may be considered a public nuisance, and that a use or interference with real property is not required. A public nuisance is one which interferes with public health and welfare and creates an imminent risk of public harm.

- 1151. Defendants, though their work with Purdue Pharma and numerous other opioid industry participants, created and continues to perpetuate and maintain a public nuisance to the citizens of the Sac and Fox Nation through the massive manufacturing and distribution of millions of doses of highly addictive, commonly abused prescription pain killers known as opioids.
- 1152. Defendants' conduct, including their misrepresentations and omissions regarding opioids, as well as efforts designed to sell as many units of controlled substances as conceivably possible, have fueled an opioid epidemic within the territorial limits of Plaintiff that constitutes a public nuisance. Defendants and their opioid clients, to include Purdue, Endo, Janssen, and Teva, knowingly exacerbated the opioid epidemic that affects entire municipalities, towns, and communities.
- 1153. Defendants' conduct, including their misrepresentations and omissions regarding opioids, as well as its efforts designed to sell as many units of controlled substances as conceivably possible, constitute unlawful acts and/or omissions of duties, that annoy, injure, or endanger the comfort, repose, health, and/or safety of others.
- 1154. As a direct and proximate result of the wrongful conduct set forth herein, Defendants negligently, intentionally, and/or unreasonably interfered with the rights of the Plaintiff to be free from unwanted injuries, addictions, diseases, sicknesses, overdoses, criminal actions, and have caused ongoing damage, harm, and inconvenience to Plaintiff and its citizens.

1155. As a direct and proximate result, Plaintiff and its citizens have been exposed to the risk of addiction to prescription opioids, have become addicted, and/or have suffered other adverse consequences from their use of the addictive prescription opioids, and have been adversely affected by the addiction and abuse of others in their communities from the highly addictive, prescription pain medication.

- 1156. The annoyance, injury, and danger to the comfort, repose, health, and safety of residents of Plaintiff includes, but is not limited to:
 - Drug overdose deaths in Oklahoma had already increased six-fold from 1999 to
 2012. In 2010, overdose deaths surpassed car crash deaths in Oklahoma.
 Defendants crafted a strategy that *tripled* OxyContin sales during that time;
 - ii. From 2004 to 2014, Oklahoma's unintentional poisoning death rate effectively doubled, 8.2 deaths per 100,000 individuals to 16.3 per 100,000. Prescription opioids contributed to the majority of those deaths. The following year, McKinsey developed and with Defendants implemented "Project Turbocharge," which was adopted as the national sales theme for the following year, under the rubric of "Evolve to Excellence";
 - iii. In 2014, during the first year of "Evolve to Excellence," Oklahoma suffered in excess of 600 deaths attributable to opioid overdoses. By 2018, the number almost doubled, to 1,132.
 - iv. Prescription opioid addiction often leads to illicit opioid use and addiction;
 - v. According to the Centers for Disease Control, past misuse of prescription opioids is the strongest risk factor for heroin initiation and use;
 - vi. Oklahoma hospitals are reporting increasing numbers of newborns testing positive for prescription medications; and

vii. Defendants crafted deceptive marketing strategies that were prepared for Purdue, purchased by Purdue, and implemented by Purdue with Defendants' ongoing assistance. These strategies enflamed, purposefully, an opioid abuse and addiction epidemic that has caused the Nation, its businesses, communities and citizens to bear enormous social and economic costs including increased health care, criminal justice, and lost work productivity expenses, among others.

1157. Defendants' conduct annoys, injures, and/or endangers the comfort, repose, health, and safety of others. In addition, Defendants' conduct caused and continues to cause harm to Plaintiff and its citizens.

1158. Plaintiff seeks to abate the public nuisance Defendants enflamed and all necessary relief to abate such public nuisance.

COUNT VI. Fraud (Actual and Constructive) and Deceit

1159. Plaintiff realleges and incorporates by reference the allegations set forth above.

1160. Defendants made and caused to be made false representations to healthcare providers working in Plaintiff's territory, and/or omitted material facts, regarding the risks, efficacy, and medical necessity of opioids, generally, and Defendants' clients' opioids, specifically. Defendants knew these representations were false, made recklessly without knowledge of the truth, and/or had no reasonable ground for believing such assertions. Specifically, Defendants knowingly and/or recklessly:

- Downplayed the substantial risks of addiction and other side-effects of opioids,
 including crafting its clients' opioid sales and marketing strategies;
- ii. Overstated the efficacy of opioids generally, including making false statements regarding the effectiveness of the drugs for treating specific subsets of the patient

population (i.e., those with osteoarthritis) and their ability to improve patient function; and

- iii. Misrepresented the medical usefulness and necessity of opioids, generally, and Purdue's and other opioid manufacturer's opioids specifically, including affirmatively marketing their drugs for off label uses (i.e., fibromyalgia osteoarthritis) without solicitation and not in response to questions from healthcare providers.
- 1161. Defendants and their clients' misrepresentations and omissions deceived others, violated public confidence, and/or injured public interests. Defendants, having chosen to craft the marketing plans used by Purdue and other opioid manufacturers to make misrepresentations to healthcare providers regarding their opioids, was under a duty to disclose the whole truth, and not disclose partial and misleading truths.
- 1162. Defendants intended healthcare providers to rely upon the false representations regarding the risks, efficacy, and medical necessity of opioids, generally, and Defendants' clients' opioids, specifically, to increase the number of opioid prescriptions made by healthcare providers.
- 1163. Healthcare providers working in Plaintiff's territory did in fact rely on the false representations made in the course of numerous opioid sales and marketing plans created by Defendants and implemented with Defendants' ongoing assistance to its clients.
- 1164. Plaintiff seeks to recover all damages caused by Defendants fraudulent representations and omissions.
- 1165. Defendants acted with knowledge and willful intent, with reckless disregard for the rights of others, and/or intentionally and with malice towards others. As such, Plaintiff seeks to recover punitive damages against Defendants.

COUNT VII: Civil Conspiracy

- 1166. Plaintiff realleges and incorporates by reference the allegations set forth above.
- 1167. Defendants and their opioid manufacturer clients, working together for decades, agreed to commit numerous unlawful acts relating to the sale and marketing of their opioid products. Defendants and their opioid clients also agreed to use unlawful means to commit lawful acts as part of these sales and marketing efforts.
- 1168. Defendants and their opioid clients agreed to pursue the unlawful act of knowingly misrepresenting the addictive nature of opioids in marketing their opioids to health care providers within Plaintiff's territory.
- 1169. Defendants and Purdue deployed the unlawful means of evading Purdue's reporting and compliance obligations to the Inspector General of the United States Department of Health and Human Services for the five years Purdue was subject to a Corporate Integrity Agreement after it pled guilty in 2007 to criminal misbranding.
- 1170. Defendants and their opioid clients discussed herein conspired to violate the Oklahoma Consumer Protection Act, 15 OK Stat § 15-753 et. seq... Defendants and their clients engaged in deceptive trade practices including, making and causing to be made misrepresentations and omissions in marketing of opioids in general, and Defendants' clients' opioids, specifically, that deceived or could reasonably be expected to deceive or mislead consumers.
- 1171. Defendants and their numerous opioid clients engaged in unfair trade practices, including, intentionally downplaying the risks, overstating the benefits, and misrepresenting the medical necessity of opioids, generally, and Defendants' clients' opioids, specifically, including for off-label uses. These practices offend established public policy and are immoral, unethical, oppressive, unscrupulous or substantially injurious to consumers.

1172. Defendants knowingly made or caused to be made false or misleading representations as to the characteristics, ingredients, uses, and benefits of opioids, generally, and Defendants' clients' opioids, specifically, by downplaying the risks of addiction and abuse, overstating the efficacy, and misrepresenting the medical necessity of opioids, generally, and Defendants' clients' opioids, specifically.

- 1173. Defendants and their numerous opioid clients agreed to deploy unlawful sales and marketing tactics to achieve the lawful purpose of maximizing ROI for Defendants' opioid clients.
- 1174. As a consequence, Defendants are jointly and severally liable with its opioid clients for the salesforce optimization and sales and marketing practices used to promote ZS' clients' opioid products, including Purdue's OxyContin, Teva's Fentora, Endo's Opana, Janssen's Nucynta, and others.
- 1175. Plaintiff was damaged as a result of the unlawful acts Defendants conspired with its clients to commit.

COUNTY VIII: Civil Aiding and Abetting

- 1176. Plaintiff realleges and incorporates by reference the allegations set forth above.
- 1177. Defendants gave substantial assistance and encouragement to Purdue and their other opioid clients regarding conduct Defendants knew to be tortious and/or in violation of a duty owed by clients to third persons, including Plaintiff.
- 1178. Plaintiff was damaged as a result of the specific conduct that Defendants encouraged and substantially assisted.

COUNT IX Unjust Enrichment

1179. Plaintiff realleges and incorporates by reference the allegations set forth above.

1180. Unjust enrichment is established where the plaintiff alleges: (a) a benefit conferred upon the defendant by the plaintiff; (b) an appreciation or knowledge by the defendant of the benefit; and (c) the acceptance or retention by the defendant of the benefit under such circumstances as to make it inequitable for the defendant to retain the benefit without the payment of its value.

- 1181. Defendants were compensated for their years of work increasing opioid sales and maximizing profits for its opioid clients.
- 1182. The compensation Defendants accepted from opioid manufacturers for maximizing sales of their deadly opioid products, by misrepresenting their addictiveness, constitutes money in the possession of Defendants that, in equity and good conscience, Defendants ought not be allowed to retain.

VII. JURY DEMAND

Plaintiff, on behalf of itself and all others similarly situated, requests a trial by jury on all issues so triable.

VIII. PRAYER FOR RELIEF

WHEREFORE, Plaintiff, on behalf of itself and all others similarly situated, respectfully prays that this Court grant the following relief:

i. Enter Judgment in favor of Plaintiff, on behalf of itself and all others similarly situated, against Defendants awarding Plaintiff its actual damages caused by the opioid epidemic, including but not limited to (1) costs for providing medical care, additional therapeutic and prescription drug purchases, and other treatments for patients suffering from opioid-related addiction or disease, including overdoses and deaths (2) costs for providing treatment, counseling and rehabilitation services, (3) costs for providing treatment of infants born with opioid-related medical conditions, (4) costs for providing care for children whose parents suffer from opioid-related disability or incapacitation, (5) costs

1		associated with law enforcement and public safety relating to the opioid epidemic, and
2		(6) costs associated with drug court and other resources expended through the judicial
3		system;
4	ii.	Order that Defendants compensate Plaintiff, on behalf of itself and all others similarly
5		
6		situated, for past and future costs to abate the ongoing public nuisance caused by the
7		opioid epidemic;
8	iii.	Order Defendants to fund an "abatement fund" for the purposes of abating the opioid
9		public nuisance;
10	iv.	Enter judgment against Defendants requiring Defendants to pay punitive damages;
11	v.	Enter judgment against Defendants awarding Plaintiff its reasonable attorneys' fees, all
12		costs and expenses, pre-judgment and post-judgment interest; and
13	vi.	All other such and further relief as this Court may deem just and proper.
14	V1.	This other such and retrief refler as this court may deem just and proper.
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17		
18	Dated: Oc	
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